

- 1 -

POLYKETIDE SYNTHASE ENZYMES AND RECOMBINANT DNA
CONSTRUCTS THEREFOR

5 Cross-Reference to Related Applications

The present application claims priority to related U.S. patent application Serial Nos. 60/102,748, filed 2 Oct. 1998; 60/139,650, filed 17 June 1999; and 60/123,810, filed 11 Mar. 1999, each of which is incorporated herein by reference.

10 Field of the Invention

The present invention relates to polyketides and the polyketide synthase (PKS) enzymes that produce them. The invention also relates generally to genes encoding PKS enzymes and to recombinant host cells containing such genes and in which expression of such genes leads to the production of polyketides. The present invention also relates to
15 compounds useful as medicaments having immunosuppressive and/or neurotrophic activity. Thus, the invention relates to the fields of chemistry, molecular biology, and agricultural, medical, and veterinary technology.

Background of the Invention

20 Polyketides are a class of compounds synthesized from 2-carbon units through a series of condensations and subsequent modifications. Polyketides occur in many types of organisms, including fungi and mycelial bacteria, in particular, the actinomycetes. Polyketides are biologically active molecules with a wide variety of structures, and the class encompasses numerous compounds with diverse activities. Tetracycline,
25 erythromycin, epothilone, FK-506, FK-520, narbomycin, picromycin, rapamycin, spinocyn, and tylosin are examples of polyketides. Given the difficulty in producing polyketide compounds by traditional chemical methodology, and the typically low production of polyketides in wild-type cells, there has been considerable interest in finding improved or alternate means to produce polyketide compounds.

- 2 -

5 This interest has resulted in the cloning, analysis, and manipulation by recombinant DNA technology of genes that encode PKS enzymes. The resulting technology allows one to manipulate a known PKS gene cluster either to produce the polyketide synthesized by that PKS at higher levels than occur in nature or in hosts that otherwise do not produce the polyketide. The technology also allows one to produce molecules that are structurally related to, but distinct from, the polyketides produced from known PKS gene clusters. See, e.g., PCT publication Nos. WO 93/13663; 95/08548; 96/40968; 97/02358; 98/27203; and 98/49315; United States Patent Nos. 4,874,748; 5,063,155; 5,098,837; 5,149,639; 5,672,491; 5,712,146; 5,830,750; and 5,843,718; and 10 Fu *et al.*, 1994, *Biochemistry* 33: 9321-9326; McDaniel *et al.*, 1993, *Science* 262: 1546-1550; and Rohr, 1995, *Angew. Chem. Int. Ed. Engl.* 34(8): 881-888, each of which is incorporated herein by reference.

15 Polyketides are synthesized in nature by PKS enzymes. These enzymes, which are complexes of multiple large proteins, are similar to the synthases that catalyze condensation of 2-carbon units in the biosynthesis of fatty acids. PKSs catalyze the biosynthesis of polyketides through repeated, decarboxylative Claisen condensations between acylthioester building blocks. The building blocks used to form complex polyketides are typically acylthioesters, such as acetyl, butyryl, propionyl, malonyl, hydroxymalonyl, methylmalonyl, and ethylmalonyl CoA. Other building blocks include 20 amino acid like acylthioesters. PKS enzymes that incorporate such building blocks include an activity that functions as an amino acid ligase (an AMP ligase) or as a non-ribosomal peptide synthetase (NRPS). Two major types of PKS enzymes are known; these differ in their composition and mode of synthesis of the polyketide synthesized. These two major types of PKS enzymes are commonly referred to as Type I or "modular" 25 and Type II "iterative" PKS enzymes.

30 In the Type I or modular PKS enzyme group, a set of separate catalytic active sites (each active site is termed a "domain", and a set thereof is termed a "module") exists for each cycle of carbon chain elongation and modification in the polyketide synthesis pathway. The typical modular PKS is composed of several large polypeptides, which can be segregated from amino to carboxy termini into a loading module, multiple extender

- 3 -

modules, and a releasing (or thioesterase) domain. The PKS enzyme known as 6-deoxyerythronolide B synthase (DEBS) is a Type I PKS. In DEBS, there is a loading module, six extender modules, and a thioesterase (TE) domain. The loading module, six extender modules, and TE of DEBS are present on three separate proteins (designated
5 DEBS-1, DEBS-2, and DEBS-3, with two extender modules per protein). Each of the DEBS polypeptides is encoded by a separate open reading frame (ORF) or gene; these genes are known as *eryAI*, *eryAII*, and *eryAIII*. See Caffrey *et al.*, 1992, *FEBS Letters* 304: 205, and U.S. Patent No. 5,824,513, each of which is incorporated herein by reference.

10 Generally, the loading module is responsible for binding the first building block used to synthesize the polyketide and transferring it to the first extender module. The loading module of DEBS consists of an acyltransferase (AT) domain and an acyl carrier protein (ACP) domain. Another type of loading module utilizes an inactivated ketosynthase (KS) domain and AT and ACP domains. This inactivated KS is in some
15 instances called KS^Q, where the superscript letter is the abbreviation for the amino acid, glutamine, that is present instead of the active site cysteine required for ketosynthase activity. In other PKS enzymes, including the FK-506 PKS, the loading module incorporates an unusual starter unit and is composed of a CoA ligase like activity domain. In any event, the loading module recognizes a particular acyl-CoA (usually acetyl or
20 propionyl but sometimes butyryl or other acyl-CoA) and transfers it as a thiol ester to the ACP of the loading module.

The AT on each of the extender modules recognizes a particular extender-CoA (malonyl or alpha-substituted malonyl, i.e., methylmalonyl, ethylmalonyl, and 2-hydroxymalonyl) and transfers it to the ACP of that extender module to form a thioester.
25 Each extender module is responsible for accepting a compound from a prior module, binding a building block, attaching the building block to the compound from the prior module, optionally performing one or more additional functions, and transferring the resulting compound to the next module.

Each extender module of a modular PKS contains a KS, AT, ACP, and zero, one,
30 two, or three domains that modify the beta-carbon of the growing polyketide chain. A

- 4 -

typical (non-loading) minimal Type I PKS extender module is exemplified by extender module three of DEBS, which contains a KS domain, an AT domain, and an ACP domain. These three domains are sufficient to activate a 2-carbon extender unit and attach it to the growing polyketide molecule. The next extender module, in turn, is responsible for attaching the next building block and transferring the growing compound to the next extender module until synthesis is complete.

Once the PKS is primed with acyl- and malonyl-ACPs, the acyl group of the loading module is transferred to form a thiol ester (trans-esterification) at the KS of the first extender module; at this stage, extender module one possesses an acyl-KS and a malonyl (or substituted malonyl) ACP. The acyl group derived from the loading module is then covalently attached to the alpha-carbon of the malonyl group to form a carbon-carbon bond, driven by concomitant decarboxylation, and generating a new acyl-ACP that has a backbone two carbons longer than the loading building block (elongation or extension).

The polyketide chain, growing by two carbons each extender module, is sequentially passed as covalently bound thiol esters from extender module to extender module, in an assembly line-like process. The carbon chain produced by this process alone would possess a ketone at every other carbon atom, producing a polyketone, from which the name polyketide arises. Most commonly, however, additional enzymatic activities modify the beta keto group of each two carbon unit just after it has been added to the growing polyketide chain but before it is transferred to the next module.

Thus, in addition to the minimal module containing KS, AT, and ACP domains necessary to form the carbon-carbon bond, and as noted above, other domains that modify the beta-carbonyl moiety can be present. Thus, modules may contain a ketoreductase (KR) domain that reduces the keto group to an alcohol. Modules may also contain a KR domain plus a dehydratase (DH) domain that dehydrates the alcohol to a double bond. Modules may also contain a KR domain, a DH domain, and an enoylreductase (ER) domain that converts the double bond product to a saturated single bond using the beta carbon as a methylene function. An extender module can also contain other enzymatic activities, such as, for example, a methylase or dimethylase activity.

After traversing the final extender module, the polyketide encounters a releasing domain that cleaves the polyketide from the PKS and typically cyclizes the polyketide. For example, final synthesis of 6-dEB is regulated by a TE domain located at the end of extender module six. In the synthesis of 6-dEB, the TE domain catalyzes cyclization of the macrolide ring by formation of an ester linkage. In FK-506, FK-520, rapamycin, and similar polyketides, the TE activity is replaced by a RapP (for rapamycin) or RapP like activity that makes a linkage incorporating a pipecolate acid residue. The enzymatic activity that catalyzes this incorporation for the rapamycin enzyme is known as RapP, encoded by the *rapP* gene. The polyketide can be modified further by tailoring enzymes; these enzymes add carbohydrate groups or methyl groups, or make other modifications, i.e., oxidation or reduction, on the polyketide core molecule. For example, 6-dEB is hydroxylated at C-6 and C-12 and glycosylated at C-3 and C-5 in the synthesis of erythromycin A.

In Type I PKS polypeptides, the order of catalytic domains is conserved. When all beta-keto processing domains are present in a module, the order of domains in that module from N-to-C-terminus is always KS, AT, DH, ER, KR, and ACP. Some or all of the beta-keto processing domains may be missing in particular modules, but the order of the domains present in a module remains the same. The order of domains within modules is believed to be important for proper folding of the PKS polypeptides into an active complex. Importantly, there is considerable flexibility in PKS enzymes, which allows for the genetic engineering of novel catalytic complexes. The engineering of these enzymes is achieved by modifying, adding, or deleting domains, or replacing them with those taken from other Type I PKS enzymes. It is also achieved by deleting, replacing, or adding entire modules with those taken from other sources. A genetically engineered PKS complex should of course have the ability to catalyze the synthesis of the product predicted from the genetic alterations made.

Alignments of the many available amino acid sequences for Type I PKS enzymes has approximately defined the boundaries of the various catalytic domains. Sequence alignments also have revealed linker regions between the catalytic domains and at the N- and C-termini of individual polypeptides. The sequences of these linker regions are less

- 6 -

well conserved than are those for the catalytic domains, which is in part how linker regions are identified. Linker regions can be important for proper association between domains and between the individual polypeptides that comprise the PKS complex. One can thus view the linkers and domains together as creating a scaffold on which the domains and modules are positioned in the correct orientation to be active. This organization and positioning, if retained, permits PKS domains of different or identical substrate specificities to be substituted (usually at the DNA level) between PKS enzymes by various available methodologies. In selecting the boundaries of, for example, an AT replacement, one can thus make the replacement so as to retain the linkers of the recipient PKS or to replace them with the linkers of the donor PKS AT domain, or, preferably, make both constructs to ensure that the correct linker regions between the KS and AT domains have been included in at least one of the engineered enzymes. Thus, there is considerable flexibility in the design of new PKS enzymes with the result that known polyketides can be produced more effectively, and novel polyketides useful as pharmaceuticals or for other purposes can be made.

By appropriate application of recombinant DNA technology, a wide variety of polyketides can be prepared in a variety of different host cells provided one has access to nucleic acid compounds that encode PKS proteins and polyketide modification enzymes. The present invention helps meet the need for such nucleic acid compounds by providing recombinant vectors that encode the FK-520 PKS enzyme and various FK-520 modification enzymes. Moreover, while the FK-506 and FK-520 polyketides have many useful activities, there remains a need for compounds with similar useful activities but with better pharmacokinetic profile and metabolism and fewer side-effects. The present invention helps meet the need for such compounds as well.

Summary of the Invention

In one embodiment, the present invention provides recombinant DNA vectors that encode all or part of the FK-520 PKS enzyme. Illustrative vectors of the invention include cosmid pKOS034-120, pKOS034-124, pKOS065-C31, pKOS065-C3, pKOS065-M27, and pKOS065-M21. The invention also provides nucleic acid compounds that

- 7 -

5 encode the various domains of the FK-520 PKS, i.e., the KS, AT, ACP, KR, DH, and ER domains. These compounds can be readily used, alone or in combination with nucleic acids encoding other FK-520 or non-FK-520 PKS domains, as intermediates in the construction of recombinant vectors that encode all or part of PKS enzymes that make novel polyketides.

10 The invention also provides isolated nucleic acids that encode all or part of one or more modules of the FK-520 PKS, each module comprising a ketosynthase activity, an acyl transferase activity, and an acyl carrier protein activity. The invention provides an isolated nucleic acid that encodes one or more open reading frames of FK-520 PKS genes, said open reading frames comprising coding sequences for a CoA ligase activity, an NRPS activity, or two or more extender modules. The invention also provides recombinant expression vectors containing these nucleic acids.

15 In another embodiment, the invention provides isolated nucleic acids that encode all or a part of a PKS that contains at least one module in which at least one of the domains in the module is a domain from a non-FK-520 PKS and at least one domain is from the FK-520 PKS. The non-FK-520 PKS domain or module originates from the rapamycin PKS, the FK-506 PKS, DEBS, or another PKS. The invention also provides recombinant expression vectors containing these nucleic acids.

20 In another embodiment, the invention provides a method of preparing a polyketide, said method comprising transforming a host cell with a recombinant DNA vector that encodes at least one module of a PKS, said module comprising at least one FK-520 PKS domain, and culturing said host cell under conditions such that said PKS is produced and catalyzes synthesis of said polyketide. In one aspect, the method is practiced with a *Streptomyces* host cell. In another aspect, the polyketide produced is FK-
25 520. In another aspect, the polyketide produced is a polyketide related in structure to FK-520. In another aspect, the polyketide produced is a polyketide related in structure to FK-506 or rapamycin.

30 In another embodiment, the invention provides a set of genes in recombinant form sufficient for the synthesis of ethylmalonyl CoA in a heterologous host cell. These genes and the methods of the invention enable one to create recombinant host cells with the

- 8 -

ability to produce polyketides or other compounds that require ethylmalonyl CoA for biosynthesis. The invention also provides recombinant nucleic acids that encode AT domains specific for ethylmalonyl CoA. Thus, the compounds of the invention can be used to produce polyketides requiring ethylmalonyl CoA in host cells that otherwise are
5 unable to produce such polyketides.

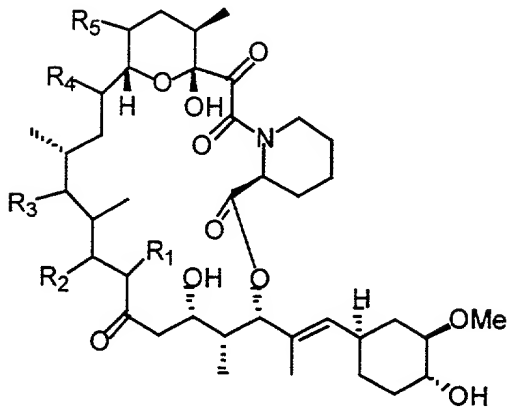
In another embodiment, the invention provides a set of genes in recombinant form sufficient for the synthesis of 2-hydroxymalonyl CoA and 2-methoxymalonyl CoA in a heterologous host cell. These genes and the methods of the invention enable one to create recombinant host cells with the ability to produce polyketides or other compounds that
10 require 2-hydroxymalonyl CoA for biosynthesis. The invention also provides recombinant nucleic acids that encode AT domains specific for 2-hydroxymalonyl CoA and 2-methoxymalonyl CoA. Thus, the compounds of the invention can be used to produce polyketides requiring 2-hydroxymalonyl CoA or 2-methoxymalonyl CoA in host cells that are otherwise unable to produce such polyketides.

In another embodiment, the invention provides a compound related in structure to FK-520 or FK-506 that is useful in the treatment of a medical condition. These compounds include compounds in which the C-13 methoxy group is replaced by a moiety selected from the group consisting of hydrogen, methyl, and ethyl moieties. Such compounds are less susceptible to the main *in vivo* pathway of degradation for FK-520
15 and FK-506 and related compounds and thus exhibit an improved pharmacokinetic profile. The compounds of the invention also include compounds in which the C-15 methoxy group is replaced by a moiety selected from the group consisting of hydrogen, methyl, and ethyl moieties. The compounds of the invention also include the above compounds further modified by chemical methodology to produce derivatives such as,
20 but not limited to, the C-18 hydroxyl derivatives, which have potent neurotrophin but not immunosuppression activities.

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- 9 -

Thus, the invention provides polyketides having the structure:



wherein, R₁ is hydrogen, methyl, ethyl, or allyl; R₂ is hydrogen or hydroxyl, provided
5 that when R₂ is hydrogen, there is a double bond between C-20 and C-19; R₃ is hydrogen
or hydroxyl; R₄ is methoxyl, hydrogen, methyl, or ethyl; and R₅ is methoxyl, hydrogen,
methyl, or ethyl; but not including FK-506, FK-520, 18-hydroxy-FK-520, and 18-
hydroxy-FK-506. The invention provides these compounds in purified form and in
pharmaceutical compositions.

10 In another embodiment, the invention provides a method for treating a medical
condition by administering a pharmaceutically efficacious dose of a compound of the
invention. The compounds of the invention may be administered to achieve
immunosuppression or to stimulate nerve growth and regeneration.

15 These and other embodiments and aspects of the invention will be more fully
understood after consideration of the attached Drawings and their brief description below,
together with the detailed description, examples, and claims that follow.

Brief Description of the Drawings

20 Figure 1 shows a diagram of the FK-520 biosynthetic gene cluster. The top line
provides a scale in kilobase pairs (kb). The second line shows a restriction map with
selected restriction enzyme recognition sequences indicated. K is *Kpn*I; X is *Xho*I, S is
*Sac*I; P is *Pst*I; and E is *Eco*RI. The third line indicates the position of FK-520 PKS and
related genes. Genes are abbreviated with a one letter designation, i.e., C is *fkbc*.

- 10 -

Immediately under the third line are numbered segments showing where the loading module (L) and ten different extender modules (numbered 1 - 10) are encoded on the various genes shown. At the bottom of the Figure, the DNA inserts of various cosmids of the invention (i.e., 34-124 is cosmid pKOS034-124) are shown in alignment with the FK-520 biosynthetic gene cluster.

Figure 2 shows the loading module (load), the ten extender modules, and the peptide synthetase domain of the FK-520 PKS, together with, on the top line, the genes that encode the various domains and modules. Also shown are the various intermediates in FK-520 biosynthesis, as well as the structure of FK-520, with carbons 13, 15, 21, and 31 numbered. The various domains of each module and subdomains of the loading module are also shown. The darkened circles showing the DH domains in modules 2, 3, and 4 indicate that the dehydratase domain is not functional as a dehydratase; this domain may affect the stereochemistry at the corresponding position in the polyketide. The substituents on the FK-520 structure that result from the action of non-PKS enzymes are also indicated by arrows, together with the types of enzymes or the genes that code for the enzymes that mediate the action. Although the methyltransferase is shown acting at the C-13 and C-15 hydroxyl groups after release of the polyketide from the PKS, the methyltransferase may act on the 2-hydroxymalonyl substrate prior to or contemporaneously with its incorporation during polyketide synthesis.

Figure 3 shows a close-up view of the left end of the FK-520 gene cluster, which contains at least ten additional genes. The ethyl side chain on carbon 21 of FK-520 (Figure 2) is derived from an ethylmalonyl CoA extender unit that is incorporated by an ethylmalonyl specific AT domain in extender module 4 of the PKS. At least four of the genes in this region code for enzymes involved in ethylmalonyl biosynthesis. The polyhydroxybutyrate depolymerase is involved in maintaining hydroxybutyryl-CoA pools during FK-520 production. Polyhydroxybutyrate accumulates during vegetative growth and disappears during stationary phase in other *Streptomyces* (Ranade and Vining, 1993, *Can. J. Microbiol.* 39:377). Open reading frames with unknown function are indicated with a question mark.

- 11 -

Figure 4 shows a biosynthetic pathway for the biosynthesis of ethylmalonyl CoA from acetoacetyl CoA consistent with the function assigned to four of the genes in the FK-520 gene cluster shown in Figure 3.

Figure 5 shows a close-up view of the right-end of the FK-520 PKS gene cluster (and of the sequences on cosmid pKOS065-C31). The genes shown include *fk bD*, *fk bM* (a methyl transferase that methylates the hydroxyl group on C-31 of FK-520), *fk bN* (a homolog of a gene described as a regulator of cholesterol oxidase and that is believed to be a transcriptional activator), *fk bQ* (a type II thioesterase, which can increase polyketide production levels), and *fk bS* (a crotonyl-CoA reductase involved in the biosynthesis of ethylmalonyl CoA).

Figure 6 shows the proposed degradative pathway for tacrolimus (FK-506) metabolism.

Figure 7 shows a schematic process for the construction of recombinant PKS genes of the invention that encode PKS enzymes that produce 13-desmethoxy FK-506 and FK-520 polyketides of the invention, as described in Example 4, below.

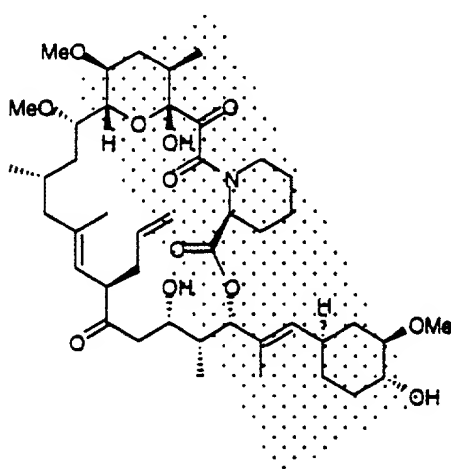
Figure 8, in Parts A and B, shows certain compounds of the invention preferred for dermal application in Part A and a synthetic route for making those compounds in Part B.

Detailed Description of the Invention

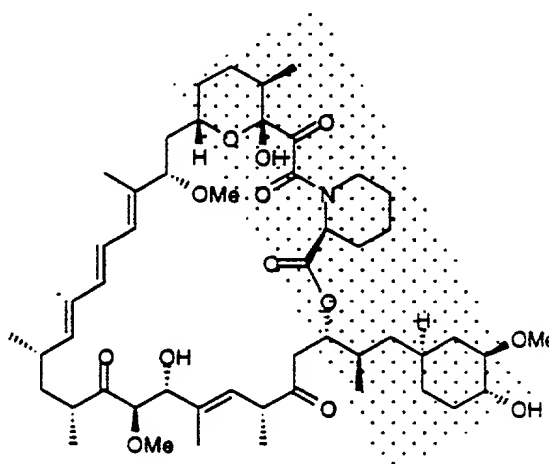
Given the valuable pharmaceutical properties of polyketides, there is a need for methods and reagents for producing large quantities of polyketides, as well as for producing related compounds not found in nature. The present invention provides such methods and reagents, with particular application to methods and reagents for producing the polyketides known as FK-520, also known as ascomycin or L-683,590 (see Holt *et al.*, 1993, *JACS* 115:9925), and FK-506, also known as tacrolimus. Tacrolimus is a macrolide immunosuppressant used to prevent or treat rejection of transplanted heart, kidney, liver, lung, pancreas, and small bowel allografts. The drug is also useful for the prevention and treatment of graft-versus-host disease in patients receiving bone marrow transplants, and for the treatment of severe, refractory uveitis. There have been additional

reports of the unapproved use of tacrolimus for other conditions, including alopecia
universalis, autoimmune chronic active hepatitis, inflammatory bowel disease, multiple
sclerosis, primary biliary cirrhosis, and scleroderma. The invention provides methods and
reagents for making novel polyketides related in structure to FK-520 and FK-506, and
5 structurally related polyketides such as rapamycin.

The FK-506 and rapamycin polyketides are potent immunosuppressants, with
chemical structures shown below.



FK-506



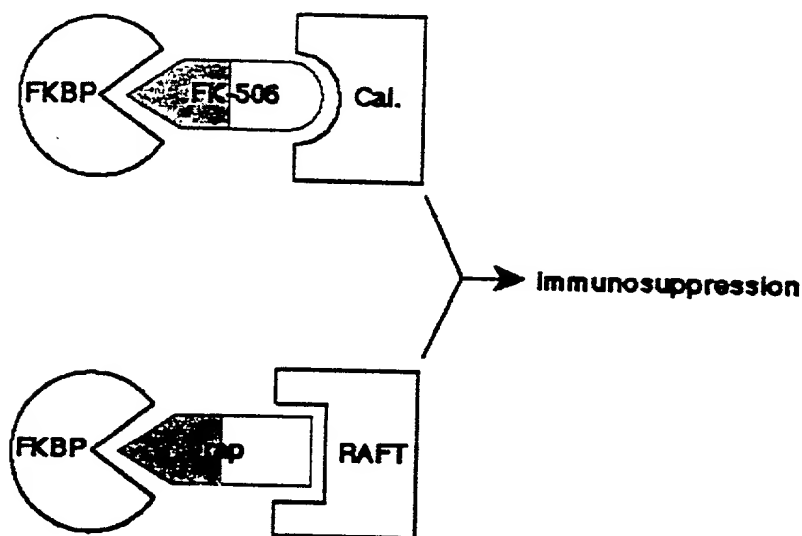
Rapamycin

FK-520 differs from FK-506 in that it lacks the allyl group at C-21 of FK-506, having
10 instead an ethyl group at that position, and has similar activity to FK-506, albeit reduced
immunosuppressive activity.

These compounds act through initial formation of an intermediate complex with
protein "immunophilins" known as FKBP (FK-506 binding proteins), including FKBP-
12. Immunophilins are a class of cytosolic proteins that form complexes with molecules
15 such as FK-506, FK-520, and rapamycin that in turn serve as ligands for other cellular
targets involved in signal transduction. Binding of FK-506, FK-520, and rapamycin to
FKBP occurs through the structurally similar segments of the polyketide molecules,
known as the "FKBP-binding domain" (as generally but not precisely indicated by the
stippled regions in the structures above). The FK-506-FKBP complex then binds
20 calcineurin, while the rapamycin-FKBP complex binds to a protein known as RAFT-1.

- 13 -

Binding of the FKBP-polyketide complex to these second proteins occurs through the dissimilar regions of the drugs known as the "effector" domains.



5 The three component FKBP-polyketide-effector complex is required for signal transduction and subsequent immunosuppressive activity of FK-506, FK-520, and rapamycin. Modifications in the effector domains of FK-506, FK-520, and rapamycin that destroy binding to the effector proteins (calcineurin or RAFT) lead to loss of immunosuppressive activity, even though FKBP binding is unaffected. Further, such analogs antagonize the immunosuppressive effects of the parent polyketides, because they compete for FKBP. Such non-immunosuppressive analogs also show reduced toxicity (see Dumont *et al.*, 1992, *Journal of Experimental Medicine* 176, 751-760), indicating that much of the toxicity of these drugs is not linked to FKBP binding.

15 In addition to immunosuppressive activity, FK-520, FK-506, and rapamycin have neurotrophic activity. In the central nervous system and in peripheral nerves, immunophilins are referred to as "neuroimmunophilins". The neuroimmunophilin FKBP is markedly enriched in the central nervous system and in peripheral nerves. Molecules that bind to the neuroimmunophilin FKBP, such as FK-506 and FK-520, have the remarkable effect of stimulating nerve growth. *In vitro*, they act as neurotrophins, i.e.,

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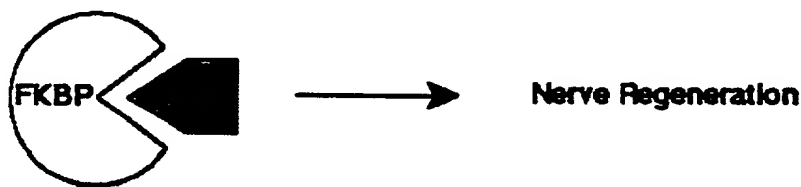
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they promote neurite outgrowth in NGF-treated PC12 cells and in sensory neuronal cultures, and in intact animals, they promote regrowth of damaged facial and sciatic nerves, and repair lesioned serotonin and dopamine neurons in the brain. See Gold *et al.*, Jun. 1999, *J. Pharm. Exp. Ther.* 289(3): 1202-1210; Lyons *et al.*, 1994, *Proc. National Academy of Science* 91: 3191-3195; Gold *et al.*, 1995, *Journal of Neuroscience* 15: 7509-7516; and Steiner *et al.*, 1997, *Proc. National Academy of Science* 94: 2019-2024.

Further, the restored central and peripheral neurons appear to be functional.

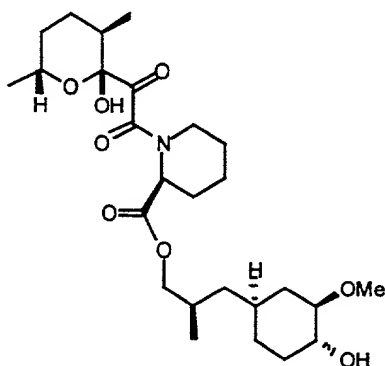
Compared to protein neurotrophic molecules (BNDF, NGF, etc.), the small-molecule neurotrophins such as FK-506, FK-520, and rapamycin have different, and often advantageous, properties. First, whereas protein neurotrophins are difficult to deliver to their intended site of action and may require intra-cranial injection, the small-molecule neurotrophins display excellent bioavailability; they are active when administered subcutaneously and orally. Second, whereas protein neurotrophins show quite specific effects, the small-molecule neurotrophins show rather broad effects. Finally, whereas protein neurotrophins often show effects on normal sensory nerves, the small-molecule neurotrophins do not induce aberrant sprouting of normal neuronal processes and seem to affect damaged nerves specifically. Neuroimmunophilin ligands have potential therapeutic utility in a variety of disorders involving nerve degeneration (e.g. multiple sclerosis, Parkinson's disease, Alzheimer's disease, stroke, traumatic spinal cord and brain injury, peripheral neuropathies).

Recent studies have shown that the immunosuppressive and neurite outgrowth activity of FK-506, FK-520, and rapamycin can be separated; the neuroregenerative activity in the absence of immunosuppressive activity is retained by agents which bind to FKBP but not to the effector proteins calcineurin or RAFT. See Steiner *et al.*, 1997, *Nature Medicine* 3: 421-428.



- 15 -

Available structure-activity data show that the important features for neurotrophic activity of rapamycin, FK-520, and FK-506 lie within the common, contiguous segments of the macrolide ring that bind to FKBP. This portion of the molecule is termed the "FKBP binding domain" (see VanDuyne *et al.*, 1993, *Journal of Molecular Biology* 229: 105-124.). Nevertheless, the effector domains of the parent macrolides contribute to conformational rigidity of the binding domain and thus indirectly contribute to FKBP binding.

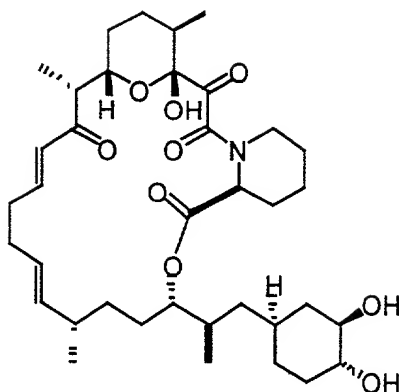


"FKBP binding domain"

There are a number of other reported analogs of FK-506, FK-520, and rapamycin that bind to FKBP but not the effector protein calcineurin or RAFT. These analogs show effects on nerve regeneration without immunosuppressive effects.

Naturally occurring FK-520 and FK-506 analogs include the antascomycins, which are FK-506-like macrolides that lack the functional groups of FK-506 that bind to calcineurin (see Fehr *et al.*, 1996, *The Journal of Antibiotics* 49: 230-233). These molecules bind FKBP as effectively as does FK-506; they antagonize the effects of both FK-506 and rapamycin, yet lack immunosuppressive activity.

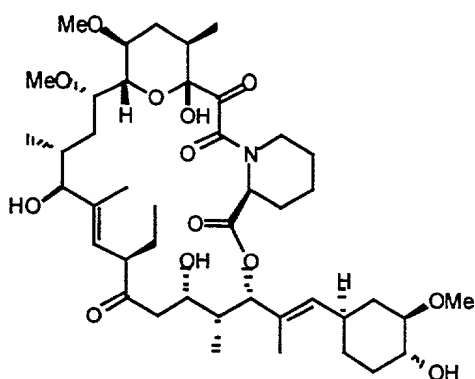
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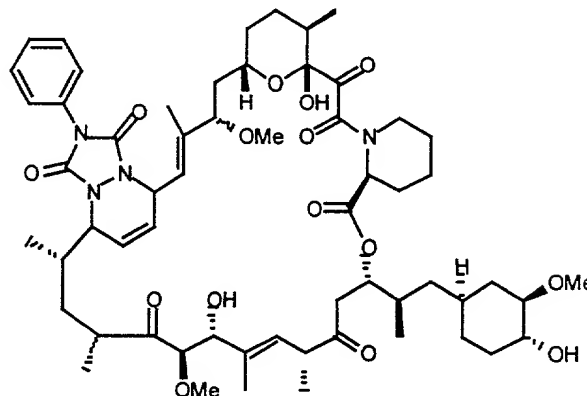
Antascomycin A

Other analogs can be produced by chemically modifying FK-506, FK-520, or rapamycin. One approach to obtaining neuroimmunophilin ligands is to destroy the effector binding region of FK-506, FK-520, or rapamycin by chemical modification.

- 5 While the chemical modifications permitted on the parent compounds are quite limited, some useful chemically modified analogs exist. The FK-520 analog L-685,818 ($ED_{50} = 0.7$ nM for FKBP binding; see Dumont *et al.*, 1992), and the rapamycin analog WAY-124,466 ($IC_{50} = 12.5$ nM; see Ocain *et al.*, 1993, *Biochemistry Biophysical Research Communications* 192: 1340-134693) are about as effective as FK-506, FK-520, and
- 10 rapamycin at promoting neurite outgrowth in sensory neurons (see Steiner *et al.*, 1997).



L-685,818

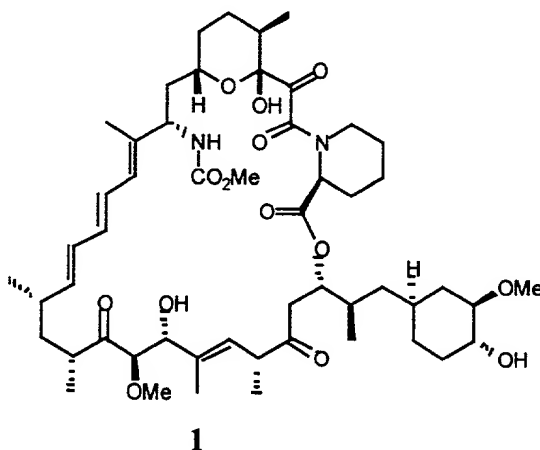


WAY-124,466

One of the few positions of rapamycin that is readily amenable to chemical modification is the allylic 16-methoxy group; this reactive group is readily exchanged by

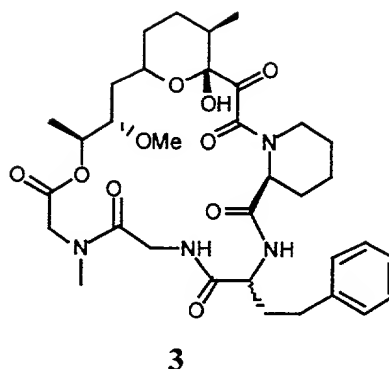
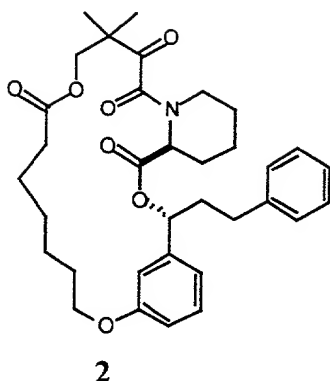
- 17 -

acid-catalyzed nucleophilic substitution. Replacement of the 16-methoxy group of rapamycin with a variety of bulky groups has produced analogs showing selective loss of immunosuppressive activity while retaining FKBP-binding (see Luengo *et al.*, 1995, *Chemistry & Biology* 2: 471-481). One of the best compounds, 1, below, shows complete
5 loss of activity in the splenocyte proliferation assay with only a 10-fold reduction in binding to FKBP.



There are also synthetic analogs of FKBP binding domains. These compounds
10 reflect an approach to obtaining neuroimmunophilin ligands based on "rationally designed" molecules that retain the FKBP-binding region in an appropriate conformation for binding to FKBP, but do not possess the effector binding regions. In one example, the ends of the FKBP binding domain were tethered by hydrocarbon chains (see Holt *et al.*, 1993, *Journal of the American Chemical Society* 115: 9925-9938); the best analog, 2,
15 below, binds to FKBP about as well as FK-506. In a similar approach, the ends of the FKBP binding domain were tethered by a tripeptide to give analog 3, below, which binds to FKBP about 20-fold poorer than FK-506. These compounds are anticipated to have neuroimmunophilin binding activity.

- 18 -



In a primate MPTP model of Parkinson's disease, administration of FKBP ligand GPI-1046 caused brain cells to regenerate and behavioral measures to improve. MPTP is a neurotoxin, which, when administered to animals, selectively damages nigral-striatal dopamine neurons in the brain, mimicking the damage caused by Parkinson's disease. Whereas, before treatment, animals were unable to use affected limbs, the FKBP ligand restored the ability of animals to feed themselves and gave improvements in measures of locomotor activity, neurological outcome, and fine motor control. There were also corresponding increases in regrowth of damaged nerve terminals. These results demonstrate the utility of FKBP ligands for treatment of diseases of the CNS.

From the above description, two general approaches towards the design of non-immunosuppressant, neuroimmunophilin ligands can be seen. The first involves the construction of constrained cyclic analogs of FK-506 in which the FKBP binding domain is fixed in a conformation optimal for binding to FKBP. The advantages of this approach are that the conformation of the analogs can be accurately modeled and predicted by computational methods, and the analogs closely resemble parent molecules that have proven pharmacological properties. A disadvantage is that the difficult chemistry limits the numbers and types of compounds that can be prepared. The second approach involves the trial and error construction of acyclic analogs of the FKBP binding domain by conventional medicinal chemistry. The advantages to this approach are that the chemistry is suitable for production of the numerous compounds needed for such interactive chemistry-bioassay approaches. The disadvantages are that the molecular types of compounds that have emerged have no known history of appropriate pharmacological

- 19 -

properties, have rather labile ester functional groups, and are too conformationally mobile to allow accurate prediction of conformational properties.

The present invention provides useful methods and reagents related to the first approach, but with significant advantages. The invention provides recombinant PKS
5 genes that produce a wide variety of polyketides that cannot otherwise be readily synthesized by chemical methodology alone. Moreover, the present invention provides polyketides that have either or both of the desired immunosuppressive and neurotrophic activities, some of which are produced only by fermentation and others of which are produced by fermentation and chemical modification. Thus, in one aspect, the invention
10 provides compounds that optimally bind to FKBP but do not bind to the effector proteins. The methods and reagents of the invention can be used to prepare numerous constrained cyclic analogs of FK-520 in which the FKBP binding domain is fixed in a conformation optimal for binding to FKBP. Such compounds will show neuroimmunophilin binding (neurotrophic) but not immunosuppressive effects. The invention also allows direct
15 manipulation of FK-520 and related chemical structures *via* genetic engineering of the enzymes involved in the biosynthesis of FK-520 (as well as related compounds, such as FK-506 and rapamycin); similar chemical modifications are simply not possible because of the complexity of the structures. The invention can also be used to introduce "chemical handles" into normally inert positions that permit subsequent chemical modifications.

20 Several general approaches to achieve the development of novel neuroimmunophilin ligands are facilitated by the methods and reagents of the present invention. One approach is to make "point mutations" of the functional groups of the parent FK-520 structure that bind to the effector molecules to eliminate their binding potential. These types of structural modifications are difficult to perform by chemical
25 modification, but can be readily accomplished with the methods and reagents of the invention.

A second, more extensive approach facilitated by the present invention is to utilize molecular modeling to predict optimal structures *ab initio* that bind to FKBP but not effector molecules. Using the available X-ray crystal structure of FK-520 (or FK-506)
30 bound to FKBP, molecular modeling can be used to predict polyketides that should

- 20 -

optimally bind to FKBP but not calcineurin. Various macrolide structures can be generated by linking the ends of the FKBP-binding domain with "all possible" polyketide chains of variable length and substitution patterns that can be prepared by genetic manipulation of the FK-520 or FK-506 PKS gene cluster in accordance with the methods of the invention. The ground state conformations of the virtual library can be determined, and compounds that possess binding domains most likely to bind well to FKBP can be prepared and tested.

Once a compound is identified in accordance with the above approaches, the invention can be used to generate a focused library of analogs around the lead candidate, to "fine tune" the compound for optimal properties. Finally, the genetic engineering methods of the invention can be directed towards producing "chemical handles" that enable medicinal chemists to modify positions of the molecule previously inert to chemical modification. This opens the path to previously prohibited chemical optimization of lead compounds by time-proven approaches.

Moreover, the present invention provides polyketide compounds and the recombinant genes for the PKS enzymes that produce the compounds that have significant advantages over FK-506 and FK-520 and their analogs. The metabolism and pharmacokinetics of tacrolimus has been extensively studied, and FK-520 is believed to be similar in these respects. Absorption of tacrolimus is rapid, variable, and incomplete from the gastrointestinal tract (Harrison's Principles of Internal Medicine, 14th edition, 1998, McGraw Hill, 14, 20, 21, 64-67). The mean bioavailability of the oral dosage form is 27%, (range 5 to 65%). The volume of distribution (V₀D) based on plasma is 5 to 65 L per kg of body weight (L/kg), and is much higher than the V₀D based on whole blood concentrations, the difference reflecting the binding of tacrolimus to red blood cells. Whole blood concentrations may be 12 to 67 times the plasma concentrations. Protein binding is high (75 to 99%), primarily to albumin and alpha₁-acid glycoprotein. The half-life for distribution is 0.9 hour; elimination is biphasic and variable: terminal-11.3 hr (range, 3.5 to 40.5 hours). The time to peak concentration is 0.5 to 4 hours after oral administration.

- 21 -

Tacrolimus is metabolized primarily by cytochrome P450 3A enzymes in the liver and small intestine. The drug is extensively metabolized with less than 1% excreted unchanged in urine. Because hepatic dysfunction decreases clearance of tacrolimus, doses have to be reduced substantially in primary graft non-function, especially in children. In addition, drugs that induce the cytochrome P450 3A enzymes reduce tacrolimus levels, while drugs that inhibit these P450s increase tacrolimus levels. Tacrolimus bioavailability doubles with co-administration of ketoconazole, a drug that inhibits P450 3A. See, Vincent *et al.*, 1992, *In vitro* metabolism of FK-506 in rat, rabbit, and human liver microsomes: Identification of a major metabolite and of cytochrome P450 3A as the major enzymes responsible for its metabolism, *Arch. Biochem. Biophys.* 294: 454-460; Iwasaki *et al.*, 1993, Isolation, identification, and biological activities of oxidative metabolites of FK-506, a potent immunosuppressive macrolide lactone, *Drug Metabolism & Disposition* 21: 971-977; Shiraga *et al.*, 1994, Metabolism of FK-506, a potent immunosuppressive agent, by cytochrome P450 3A enzymes in rat, dog, and human liver microsomes, *Biochem. Pharmacol.* 47: 727-735; and Iwasaki *et al.*, 1995, Further metabolism of FK-506 (Tacrolimus); Identification and biological activities of the metabolites oxidized at multiple sites of FK-506, *Drug Metabolism & Disposition* 23: 28-34. The cytochrome P450 3A subfamily of isozymes has been implicated as important in this degradative process.

Structures of the eight isolated metabolites formed by liver microsomes are shown in Figure 6. Four metabolites of FK-506 involve demethylation of the oxygens on carbons 13, 15, and 31, and hydroxylation of carbon 12. The 13-demethylated (hydroxy) compounds undergo cyclizations of the 13-hydroxy at C-10 to give MI, MVI and MVII, and the 12-hydroxy metabolite at C-10 to give I. Another four metabolites formed by oxidation of the four metabolites mentioned above were isolated by liver microsomes from dexamethasone treated rats. Three of these are metabolites doubly demethylated at the methoxy groups on carbons 15 and 31 (M-V), 13 and 31 (M-VI), and 13 and 15 (M-VII). The fourth, M-VIII, was the metabolite produced after demethylation of the 31-methoxy group, followed by formation of a fused ring system by further oxidation.

Among the eight metabolites, M-II has immunosuppressive activity comparable to that of

- 22 -

FK-506, whereas the other metabolites exhibit weak or negligible activities. Importantly, the major metabolite of human, dog, and rat liver microsomes is the 13-demethylated and cyclized FK-506 (M-I).

Thus, the major metabolism of FK-506 proceeds via 13-demethylation followed
5 by cyclization to the inactive M-I, this representing about 90% of the metabolic products after a 10 minute incubation with liver microsomes. Analogs of tacrolimus that do not possess a C-13 methoxy group would not be susceptible to the first and most important biotransformation in the destructive metabolism of tacrolimus (i.e. cyclization of 13-hydroxy to C-10). Thus, a 13-desmethoxy analog of FK-506 should have a longer half-
10 life in the body than does FK-506. The C-13 methoxy group is believed not to be required for binding to FKBP or calcineurin. The C-13 methoxy is not present on the identical position of rapamycin, which binds to FKBP with equipotent affinity as tacrolimus. Also, analysis of the 3-dimensional structure of the FKBP-tacrolimus-calcineurin complex shows that the C-13 methoxy has no interaction with FKBP and only
15 a minor interaction with calcineurin. The present invention provides C-13-desmethoxy analogs of FK-506 and FK-520, as well as the recombinant genes that encode the PKS enzymes that catalyze their synthesis and host cells that produce the compounds.

These compounds exhibit, relative to their naturally occurring counterparts, prolonged immunosuppressive action *in vivo*, thereby allowing a lower dosage and/or
20 reduced frequency of administration. Dosing is more predictable, because the variability in FK-506 dosage is largely due to variation of metabolism rate. FK-506 levels in blood can vary widely depending on interactions with drugs that induce or inhibit cytochrome P450 3A (summarized in USP Drug Information for the Health Care Professional). Of particular importance are the numerous drugs that inhibit or compete for CYP 3A,
25 because they increase FK-506 blood levels and lead to toxicity (Prograf package insert, Fujisawa □ US, Rev 4/97, Rec 6/97). Also important are the drugs that induce P450 3A (e.g. Dexamethasone), because they decrease FK-506 blood levels and reduce efficacy. Because the major site of CYP 3A action on FK-506 is removed in the analogs provided by the present invention, those analogs are not as susceptible to drug interactions as the
30 naturally occurring compounds.

- 23 -

Hyperglycemia, nephrotoxicity, and neurotoxicity are the most significant adverse effects resulting from the use of FK-506 and are believed to be similar for FK-520.

Because these effects appear to occur primarily by the same mechanism as the immunosuppressive action (i.e. FKBP-calcineurin interaction), the intrinsic toxicity of the desmethoxy analogs may be similar to FK-506. However, toxicity of FK-506 is dose related and correlates with high blood levels of the drug (Prograf package insert, Fujisawa □ US, Rev 4/97, Rec 6/97). Because the levels of the compounds provided by the present invention should be more controllable, the incidence of toxicity should be significantly decreased with the 13-desmethoxy analogs. Some reports show that certain FK-506 metabolites are more toxic than FK-506 itself, and this provides an additional reason to expect that a CYP 3A resistant analog can have lower toxicity and a higher therapeutic index.

Thus, the present invention provides novel compounds related in structure to FK-506 and FK-520 but with improved properties. The invention also provides methods for making these compounds by fermentation of recombinant host cells, as well as the recombinant host cells, the recombinant vectors in those host cells, and the recombinant proteins encoded by those vectors. The present invention also provides other valuable materials useful in the construction of these recombinant vectors that have many other important applications as well. In particular, the present invention provides the FK-520 PKS genes, as well as certain genes involved in the biosynthesis of FK-520 in recombinant form.

FK-520 is produced at relatively low levels in the naturally occurring cells, *Streptomyces hygroscopicus* var. *ascomyceticus*, in which it was first identified. Thus, another benefit provided by the recombinant FK-520 PKS and related genes of the present invention is the ability to produce FK-520 in greater quantities in the recombinant host cells provided by the invention. The invention also provides methods for making novel FK-520 analogs, in addition to the desmethoxy analogs described above, and derivatives in recombinant host cells of any origin.

The biosynthesis of FK-520 involves the action of several enzymes. The FK-520 PKS enzyme, which is composed of the *fkfA*, *fkfB*, *fkfC*, and *fkfP* gene products,

- 24 -

synthesizes the core structure of the molecule. There is also a hydroxylation at C-9 mediated by the P450 hydroxylase that is the *fk bD* gene product and that is oxidized by the *fk bO* gene product to result in the formation of a keto group at C-9. There is also a methylation at C-31 that is mediated by an O-methyltransferase that is the *fk bM* gene product. There are also methylations at the C-13 and C-15 positions by a methyltransferase believed to be encoded by the *fk bG* gene; this methyltransferase may act on the hydroxymalonyl CoA substrates prior to binding of the substrate to the AT domains of the PKS during polyketide synthesis. The present invention provides the genes encoding these enzymes in recombinant form. The invention also provides the genes encoding the enzymes involved in ethylmalonyl CoA and 2-hydroxymalonyl CoA biosynthesis in recombinant form. Moreover, the invention provides *Streptomyces hygroscopicus* var. *asco myceticus* recombinant host cells lacking one or more of these genes that are useful in the production of useful compounds.

The cells are useful in production in a variety of ways. First, certain cells make a useful FK-520-related compound merely as a result of inactivation of one or more of the FK-520 biosynthesis genes. Thus, by inactivating the C-31 O-methyltransferase gene in *Streptomyces hygroscopicus* var. *asco myceticus*, one creates a host cell that makes a desmethyl (at C-31) derivative of FK-520. Second, other cells of the invention are unable to make FK-520 or FK-520 related compounds due to an inactivation of one or more of the PKS genes. These cells are useful in the production of other polyketides produced by PKS enzymes that are encoded on recombinant expression vectors and introduced into the host cell.

Moreover, if only one PKS gene is inactivated, the ability to produce FK-520 or an FK-520 derivative compound is restored by introduction of a recombinant expression vector that contains the functional gene in a modified or unmodified form. The introduced gene produces a gene product that, together with the other endogenous and functional gene products, produces the desired compound. This methodology enables one to produce FK-520 derivative compounds without requiring that all of the genes for the PKS enzyme be present on one or more expression vectors. Additional applications and benefits of such cells and methodology will be readily apparent to those of skill in the art

- 25 -

after consideration of how the recombinant genes were isolated and employed in the construction of the compounds of the invention.

The FK-520 biosynthetic genes were isolated by the following procedure. Genomic DNA was isolated from *Streptomyces hygroscopicus* var. *ascomyceticus* (ATCC 14891) using the lysozyme/proteinase K protocol described in Genetic Manipulation of *Streptomyces* - A Laboratory Manual (Hopwood *et al.*, 1986). The average size of the DNA was estimated to be between 80 - 120 kb by electrophoresis on 0.3% agarose gels. A library was constructed in the SuperCos™ vector according to the manufacturer's instructions and with the reagents provided in the commercially available kit (Stratagene). Briefly, 100 µg of genomic DNA was partially digested with 4 units of *Sau*3A I for 20 min. in a reaction volume of 1 mL, and the fragments were dephosphorylated and ligated to SuperCos vector arms. The ligated DNA was packaged and used to infect log-stage XL1-BlueMR cells. A library of about 10,000 independent cosmid clones was obtained.

Based on recently published sequence from the FK-506 cluster (Motamedi and Shafiee, 1998, *Eur. J. Biochem.* 256: 528), a probe for the *fkfO* gene was isolated from ATCC 14891 using PCR with degenerate primers. With this probe, a cosmid designated pKOS034-124 was isolated from the library. With probes made from the ends of cosmid pKOS034-124, an additional cosmid designated pKOS034-120 was isolated. These cosmids (pKOS034-124 and pKOS034-120) were shown to contain DNA inserts that overlap with one another. Initial sequence data from these two cosmids generated sequences similar to sequences from the FK-506 and rapamycin clusters, indicating that the inserts were from the FK-520 PKS gene cluster. Two *Eco*RI fragments were subcloned from cosmids pKOS034-124 and pKOS034-120. These subclones were used to prepare shotgun libraries by partial digestion with *Sau*3AI, gel purification of fragments between 1.5 kb and 3 kb in size, and ligation into the pLitmus28 vector (New England Biolabs). These libraries were sequenced using dye terminators on a Beckmann CEQ2000 capillary electrophoresis sequencer, according to the manufacturer's protocols.

To obtain cosmids containing sequence on the left and right sides of the sequenced region described above, a new cosmid library of ATCC 14891 DNA was

- 26 -

prepared essentially as described above. This new library was screened with a new *fk bM* probe isolated using DNA from ATCC 14891. A probe representing the *fk bP* gene at the end of cosmid pKOS034-124 was also used. Several additional cosmids to the right of the previously sequenced region were identified. Cosmids pKOS065-C31 and pKOS065-C3
5 were identified and then mapped with restriction enzymes. Initial sequences from these cosmids were consistent with the expected organization of the cluster in this region. More extensive sequencing showed that both cosmids contained in addition to the desired sequences, other sequences not contiguous to the desired sequences on the host cell chromosomal DNA. Probing of additional cosmid libraries identified two additional
10 cosmids, pKOS065-M27 and pKOS065-M21, that contained the desired sequences in a contiguous segment of chromosomal DNA. Cosmids pKOS034-124, pKOS034-120, pKOS065-M27, and pKOS065-M21 have been deposited with the American Type Culture Collection, Manassas, VA, USA. The complete nucleotide sequence of the coding sequences of the genes that encode the proteins of the FK-520 PKS are shown
15 below but can also be determined from the cosmids of the invention deposited with the ATCC using standard methodology.

Referring to Figures 1 and 3, the FK-520 PKS gene cluster is composed of four open reading frames designated *fk bB*, *fk bC*, *fk bA*, and *fk bP*. The *fk bB* open reading frame encodes the loading module and the first four extender modules of the PKS. The *fk bC*
20 open reading frame encodes extender modules five and six of the PKS. The *fk bA* open reading frame encodes extender modules seven, eight, nine, and ten of the PKS. The *fk bP* open reading frame encodes the NRPS of the PKS. Each of these genes can be isolated from the cosmids of the invention described above. The DNA sequences of these genes are provided below preceded by the following table identifying the start and stop codons
25 of the open reading frames of each gene and the modules and domains contained therein.

<u>Nucleotides</u>	<u>Gene or Domain</u>
complement (412 - 1836)	<i>fk bW</i>
complement (2020 - 3579)	<i>fk bV</i>
complement (3969 - 4496)	<i>fk bR2</i>
complement (4595 - 5488)	<i>fk bR1</i>
5601 - 6818	<i>fk bE</i>

- 27 -

	6808 - 8052	<i>fkfF</i>
	8156 - 8824	<i>fkfG</i>
	complement (9122 - 9883)	<i>fkfH</i>
	complement (9894 - 10994)	<i>fkfI</i>
5	complement (10987 - 11247)	<i>fkfJ</i>
	complement (11244 - 12092)	<i>fkfK</i>
	complement (12113 - 13150)	<i>fkfL</i>
	complement (13212 - 23988)	<i>fkfC</i>
	complement (23992 - 46573)	<i>fkfB</i>
10	46754 - 47788	<i>fkfO</i>
	47785 - 52272	<i>fkfP</i>
	52275 - 71465	<i>fkfA</i>
	71462 - 72628	<i>fkfD</i>
	72625 - 73407	<i>fkfM</i>
15	complement (73460 - 76202)	<i>fkfN</i>
	complement (76336 - 77080)	<i>fkfQ</i>
	complement (77076 - 77535)	<i>fkfS</i>
	complement (44974 - 46573)	CoA ligase of loading domain
	complement (43777 - 44629)	ER of loading domain
20	complement (43144 - 43660)	ACP of loading domain
	complement (41842 - 43093)	KS of extender module 1 (KS1)
	complement (40609 - 41842)	AT1
	complement (39442 - 40609)	DH1
	complement (38677 - 39307)	KR1
25	complement (38371 - 38581)	ACP1
	complement (37145 - 38296)	KS2
	complement (35749 - 37144)	AT2
	complement (34606 - 35749)	DH2 (inactive)
	complement (33823 - 34480)	KR2
30	complement (33505 - 33715)	ACP2
	complement (32185 - 33439)	KS3
	complement (31018 - 32185)	AT3
	complement (29869 - 31018)	DH3 (inactive)
	complement (29092 - 29740)	KR3
35	complement (28750 - 28960)	ACP3
	complement (27430 - 28684)	KS4
	complement (26146 - 27430)	AT4
	complement (24997 - 26146)	DH4 (inactive)
	complement (24163 - 24373)	ACP4
40	complement (22653 - 23892)	KS5
	complement (21420 - 22653)	AT5
	complement (20241 - 21420)	DH5
	complement (19464 - 20097)	KR5
	complement (19116 - 19326)	ACP5

- 28 -

	complement (17820 - 19053)	KS6
	complement (16587 - 17820)	AT6
	complement (15438 - 16587)	DH6
	complement (14517 - 15294)	ER6
5	complement (13761 - 14394)	KR6
	complement (13452 - 13662)	ACP6
	52362 - 53576	KS7
	53577 - 54716	AT7
	54717 - 55871	DH7
10	56019 - 56819	ER7
	56943 - 57575	KR7
	57710 - 57920	ACP7
	57990 - 59243	KS8
	59244 - 60398	AT8
15	60399 - 61412	DH8 (inactive)
	61548 - 62180	KR8
	62328 - 62537	ACP8
	62598 - 63854	KS9
	63855 - 65084	AT9
20	65085 - 66254	DH9
	66399 - 67175	ER9
	67299 - 67931	KR9
	68094 - 68303	ACP9
	68397 - 69653	KS10
25	69654 - 70985	AT10
	71064 - 71273	ACP10

	1	GATCTCAGGC	ATGAAGTCCT	CCAGGCGAGG	CGCCGAGGTG	GTGAACACCT	CGCCGCTGCT
30	61	TGTACGGACC	ACTTCAGTCA	GCGGCGATTG	CGGAACCAAG	TCATCCGGAA	TAAAGGGCGG
	121	TTACAAGATC	CTCACATTGC	GCGACCGCCA	GCATACGCTG	AGTTGCCTCA	GAGGCAAACC
	181	GAAAGGGCGC	GGGCGGTCCG	CACCAGGGCG	GAGTACGCGA	CGAGAGTGGC	GCACCCGCGC
	241	ACCGTCACCT	CTCTCCCCCG	CCGGCGGGAT	GCCC GGCGTG	ACACGGTTGG	GCTCTCCTCG
	301	ACGCTGAACA	CCCGCGCGGT	GTGGCGTCGG	GGACACCGCC	TGGCATCGGC	CGGGTGACGG
35	361	TACGGGGAGG	GCGTACGGCG	GCCGTGGCTC	GTGCTCACGG	CCGCCGGGCG	GTCATCCGTC
	421	GAGACGGCAC	TCGGCGAGCA	GGGACGCCTG	GTCGGCACCT	GCGGGCCGGA	CGACCGTGTG
	481	GTTCGCGGGC	GGGCGGTGGC	CGGTGGTGAG	CCAGCTCTCC	AGGGCGGTGA	AGGCTGAGCG
	541	GTGACACGGC	AGCAAAGGCC	GGAGTCGGTC	GGGGAAGGTG	TCGACGAGGG	CGTCGGTGTG
	601	CGTGCCGTCC	TCGATGCGGT	AGTAGCGGTA	CCGGCCGCCA	GGCCGCTGCC	GGACATACGC
40	661	GCGTACACGT	CGGAGCCCGG	GCGGCAGGCA	GCAGCACGTC	GAGAGTGCCCT	GGATGGTGAT
	721	CAGCGGCTTG	CCGATACGAC	CGGTCAACGC	GATGCGTTCC	ACGGCCGCGT	GGACGCCGGA
	781	GGAGCGGGTG	GCGTAGTCGT	AGTCGGCATC	GCAGCCCGGG	ACCGTCCCCG	GGTAGACAATA
	841	CGGTGTGCCG	GCTTCCTTCT	CCCCATCGAA	GCCGGGGTCG	AACTCCTCGC	GGTAGACGCG
	901	CTGCGTCAGA	TCCAGTAGA	CCTCGTGGTG	GTACGGCCAC	AAGAACTCGG	AGTCGGCCCG
45	961	GAACCCGGCG	CGGAGCAGCG	CCTCGCGCGC	CTGGCCGGCT	GCGGGGCCCG	CTGCCGCGTA
	1021	GGTGGGGTAG	TCGCGCAGGG	CGGCCGGCAG	GAAGGTGAAG	AGGTTGGGAC	CCTCCGCGCG
	1081	CCACAGGGTG	CCTTCCCAGT	CGACTCCTCC	GTCGTACAGC	TCGGGATGGT	TCTCCAGCTG
	1141	CCAGCGCACG	AGGTAGCCGC	CGTTGGACAT	CCCGGTGACC	AGGGTGCGCT	CGAGCGGCCG
	1201	GTGGTAGCGC	TGGGCGACCG	ACGCGCGGGC	GGCCCGGGTC	AGCTGGGTGA	GGCGGGTGT

- 29 -

1261	CCACTCGGCG	ACGGCGTCGC	CCGGCCGGGA	GCCATCACGG	TAGAACGCGG	GGCCGGTGT
1321	GCCCTTGTCG	GTGGCGGCGT	AGGCGTAACC	GCGGGCGAGC	ACCCAGTCGG	CGATGGCCCC
1381	GTCGTTGGCG	TACTGCTCGC	GGTTACCGGG	GGTGCCGGCC	ACGACCAGGC	CACCGTTCCA
1441	GCGGTCGGGC	AGCCGGATGA	CGAACTGGGC	GTCGTGGTTC	CACCCGTGGT	TGGTGTGGT
5	1501	GGTGGAGGTG	TCGGGGAAGT	AGCCGTCGAT	CTGGATCCCG	GGCACTCCGG
	1561	CAGGTTCTTG	GGCGTCAGCC	CTGCCCAGTC	CGCCGGGTCG	GTGTGGCCCG
	1621	TCCCGCCGTG	GTCAGCTCGT	CCAGGCAGTC	GGCCTGCTGA	CGTGCCGCCG
	1681	CAGCTGGGAC	AGACGGGCGC	AGTGACCGTC	CGGGGCATCG	GGAGCAGGCC
	1741	CGGTGAGGGG	AGCAGGACGG	CGACTGCGGC	CAGGGTGAGA	GCGCCGAGGC
10	1801	TCTCGGGGCC	CGTCCGACAC	CGAGGGGCAG	AACCATGGAG	AGCCTCCAGA
	1861	GATGACGGAC	TGGAGGCTAG	GTGCGGCACG	GTGGAGACGA	ACATGGGTGC
	1921	ACTGAGGCCC	CTCAGAGGTG	GGCCGCCGCC	ATGACGGGCG	CGGGACCGCG
	1981	GGCGGTGCCC	GCGGCCGCCA	CCGGTTCCGG	GTCCCGGGGT	CAGGGACAGG
	2041	GACGGTGAAG	TAGCCGGTCG	GCGACTCTTT	CAAGGTGGTC	GTGACGAAGG
15	2101	GCCCATGTTT	TGGCCGGAGC	CCTTGCGTA	GGTGTAAACG	GCGCTCGTCG
	2161	CGCCTGGACG	TGAGCGTAGT	TGCCGGCGGT	CCAGCAGACG	GCCGTGGCAC
	2221	CGCGGTGACC	GCGCCCAGAG	GCGGTCCGGC	CTTGCCGTCC	GCGTCCCGGG
	2281	GTAGGTGTGC	GATGTGCCCG	CCCTCAGGCC	GGTGTCCGTG	TACGACGTCG
	2341	GGTGATCTGG	GCACCGTCGC	GGTGGACGGC	GTAGTCGGTG	GCGCCGTCGA
20	2401	GGTCAGGCTG	ATGGTGGTGT	CGGTGGCGCC	GGTGGCGGCC	AGGCCGGACG
	2461	CGAACCAGGG	TCGGAGGCGG	ATCCGCTCAG	GCCGAAGAAC	TGCGTGATCC
	2521	ACAGATCGAG	TCCAGGAAGT	AGGCGGCGCC	GGTGCTGCCG	CACTGCTGTG
	2581	GGGATCGACC	GGGGTGCCGT	GCCCGATGCC	CGGCACCCCG	TTCACCTCCA
25	2641	TCCGTCCGCG	GCCAGGTACT	CCTCGTGCCG	GGTGGAGTTC	GGGCCGATCA
	2701	GTCCGGCGTC	TGGGACACGC	CGTGACACAG	GGTCCACTGG	TCGCGCAACT
	2761	GCGCGGCGCG	ACGGTGGTGT	CCTTGTCGCC	GTGCCAGATG	GCCACGCGCG
	2821	CGACCACGAG	GGGTAGCCGT	CACGGACCCG	CCGCGCCAC	TGGTCCGCGG
	2881	CCCGGGGTTC	ATGCACAGGT	ACGCGCTGCT	GACGTCGGTG	GCACAGCCGA
30	2941	GGCGACGACC	GCGCCGGCCT	GGAAGAGGTC	CGGATAGGTG	GCGAGCATCA
	3001	GGCACCGCCG	GCGGACAGCC	CGGTGATGTA	GGTGCGCTGG	GGGTCCGCGC
	3061	GACGGTGTGA	GCGGCCATCT	GCCGGATCGA	CGCGGCTTCG	CCCTGGCCCC
	3121	GCTGCTCTGG	AACCAAGTTGA	AGCACCTGTT	CGCGTTGTTC	GACGACGTGG
	3181	CACGAGCAGG	AAGCCATAGC	GGTCCGCGAA	TGAGAGCAGG	CCGGAGTTGT
35	3241	CTGGGCGTCC	TGGGTGCAAC	CGTGACAGGC	GAACACCACC	GCCGGCTCCG
	3301	CGCGGGCCCG	TAGACGTACA	TGTTACAGCCG	GCCCGGGTTC	GTGCCGAAGT
	3361	GGTCAGGTCC	GCCTTGGTCA	GACCGGGCTT	GGCCAGGCC	GCCGCGGCGT
	3421	CGCCGGGCGG	AGCAGGGCCG	CTCCGAGTAC	GAGGGCCACG	ACGGCCACGA
	3481	CACCCCCCGC	CGTCCCGGAC	GCGACAACGA	CCCGACCGGC	GGCGAGGAGG
	3541	CAGCGGGGTG	AGGATTCCCC	GGAACGGCGG	CGGCTGCATG	GCGGCTCCCT
40	3601	GGGGGGACAC	GGAGGGGTCC	CTGACGTCGA	TCAGTGGGAG	CGCCCCGGTG
	3661	TAGGGGTGGT	TCAACCCGCA	ACGGTATGGC	CCGGAGCACC	ACACCCCGCA
	3721	TGCGCCCGGA	CGGATTGTGT	CGCCTTGCGG	AATCTGATAC	CCGGACGCGA
	3781	ACCCGACACG	GGTAGGGCGT	CATGGTGTCC	GACTCGGCCG	GTCGGCCTTG
45	3841	ACGGACCGGG	CGTCGGCGGA	CCGGGCGTCG	GCGGGCTGGG	CGGTATGGCG
	3901	CCAGCCGCGT	GGGGCGGCCG	CGCCCAAGTG	CAGTACGCCG	ACCGTGGCCG
	3961	CGGACCGGTC	AGTGCAATCC	CGCGGCCCTG	CGGGACCGCT	CGTCCCAGAG
	4021	GCGGCGAACC	GGGGTCCGTG	TCCGCGCGTG	TAGACCATCA	GTGTCCGCTC
	4081	ACGATGACAC	CGTCTGGTGT	GTAGCCGATG	GTGCGCACGC	TGATGATGCC
	4141	CGGCTGGCGG	ACTCCCGGGT	GTTACAGGACC	TCGGACTGCG	AGTAGATGGT
50	4201	AAGACCGGGT	TCGGCAGCCT	GACCCGGTCC	CAGCCGAGGT	TGGCCATCAC
	4261	ATGTCGGTGA	CGCTCTGCCC	GGTGACCAGG	GCGAGGGTGA	AGGTGGAGTC
	4321	TTGCCCCAGG	TGGTGCCCGC	CGAGTAGTGG	CGGTGCAAGT	GCAGCGGCGC
	4381	GTCAGGAGCG	TGAGCCAGGA	GTTGTCCGTC	TCCAGGACCG	TGCGGCCAG
	4441	TACACGTCGC	CGGTGGTGAA	GTCCTCGAAG	TAGCGGCCCT	GCCAGCCCTC

- 30 -

5	4501	GTGCGGGTGG	CGTCCTGGTC	CGGGTTCTCA	GTCGTCATGG	CGCTCATTCT	GGGAAGTCCC
	4561	CGGTCCGCTG	TGAAATGCCG	AACCTTCACC	GGGCTCATA	GTGCGGCGCA	TGAGCCCTGG
	4621	ACCGTACGTA	GTCGTAGAAC	CTCGCCACCA	CTGGCGCGCG	TGGTCTCTCC	GCGAGTGTGA
	4681	CCACGCCGAC	CGTGCGCCGC	GCCTGCGGGT	CGTCGAGCGG	CACGGCGACG	GCGTGGTCAC
	4741	CGGGCCCCGA	CGGGCTGCCG	GTGAGGGGGG	CGACGGCCAC	ACCGAGGCCG	GCGGCGACCA
	4801	GGGCCCCGAG	CGTGCTCAGC	TCGGTGCTCT	CCAGGACGAC	CCGCGGCACG	AATCCGGCCG
	4861	CGGCGCACAG	CCGGTCGGTG	ATCTGGCGCA	GTCCGAAGAC	CGGCTCCAGT	GCCACGAACG
	4921	CCTCATCGGC	CAGCTCCGCG	GTCCGCACCC	GGCGGCGTCT	GGCCAGCCCG	TGTCCGGGTG
10	4981	GGACGAGCAG	GCACAGTGCC	TCGTCCC GCA	GTGGTGTCCA	CTCCACATCG	TCCCCGGCGG
	5041	GTGCTGGGCT	GGTCAGCCCC	AGGTCCAGCC	TGCTGTTGCG	GACGTCGTCG	ACCACGGCGT
	5101	CGGCGCGGTC	GCCGCGCAGT	TCGAAGGTGG	TGCCGGGAGC	CAGCCGGCGG	TACCCGGCGA
	5161	GGAGGTCGGG	CACCAAGCCG	GTGCCGTAGG	AGTGCAGGAA	ACCCAGTGCC	ACGGTGCCCG
	5221	TGTCGGGGTC	GATCAGGGCG	GTGATGCGCT	GCTCGGCGCC	GGAGACCTCA	CTGATCGCGC
	5281	GCAGGGCGTG	GGCGCGGAAG	ACCTCGCCGT	ACTTGTGAG	CCGGAGCCCG	TTCTGGTGCC
15	5341	GGTCGAACAG	CGGCACGCCC	ACTCGTCGCT	CCAGCCGCCC	GATGGCCCTG	GACAGGGTCG
	5401	GCTGGGAGAT	GTTGAGCCGT	TCCGCGGTGA	TCGTACGCTG	CTCGTGCTCG	GCCAAAGGCCG
	5461	TGAACCACTG	CAACTCCCGT	ATCTCCATGC	AGGGACTATA	CGTACCGGGC	ATGGTCTCTG
	5521	CGAGGTTTCG	TCATTTTACA	GCGGCCGGGC	GGCGGCCCCAC	AGTGAGTCCT	CACCAACCAG
20	5581	GACCCCATGG	GAGGGACCCC	ATGTCCGAGC	CGCATCCTCG	CCCTGAACAG	GAACGCCCCG
	5641	CCGGGCCCCCT	GTCCGGTCTG	CTCGTGGTTT	CTTTGGAGCA	GGCCGTCGCC	GCTCCGTTTCG
	5701	CCACCCGCCA	CCTGGCGGAC	CTGGGCGCCC	GTGTCATCAA	GATCGAACGC	CCCGGCACGC
	5761	GCGACCTCGC	CCGCGGCTAC	GACCGCACGG	TGCGTGGCAT	GTCCAGCCAC	TTCTGCTGGC
	5821	TGAACCGGGG	GAAGGAGAGC	GTCCAGCTCG	ATGTGCGCTC	GCCGGAGGGC	AACCGGCACC
25	5881	TGCACGCCTT	GGTGGACCGG	GCCGATGTCC	TGGTGAGAA	TCTGGCACCC	GCGCCGCGG
	5941	GCCGCCTGGC	ATCGGCCACC	AGGTCTCTCG	GCGGAGCCAC	CGAGGCTGAT	CACCTGCGGA
	6001	CATATCCGGC	TACGGCAGTA	CCGGCTGCTA	CCGCGGACCG	CAAGGCGTAC	GACCTCCTTG
	6061	TCCAGTGCGA	AGCGGGGCTG	GTCTCCATCA	CCGGCACCCC	CGAGACCCCG	TCCAAGGTGG
	6121	GCCTGTCCAT	CGCGGACATC	TGTGCGGGGA	TGTACGCGTA	CTCCGGCATC	CTCACGGCCC
30	6181	TGCTGAAGCG	GGCCCGCACC	GGCCGGGGCT	CGCAGTTGGA	GGTCTCGATG	CTCGAAGCCC
	6241	TCGGTGAATG	GATGGGATAC	GCCGAGTACT	ACACGCGCTA	CGGCGGCACC	GCTCCGGCCC
	6301	GCGCCGGCGC	CAGCCACGCG	ACGATCGCCC	CCTACGGCCC	GTTACACACG	CGCGACGGGC
	6361	AGACGATCAA	TCTCGGGCTC	CAGAACGAGC	GGGAGTGGGC	TTCTTCTGTC	GGTGTCTGTC
	6421	TACAACGCCC	CGGTCTCTGC	GACGACCCGC	GCTTTTCCGG	CAACGCCGAC	CGGGTGGCGC
35	6481	ACCGCACCGA	GCTCGACGCC	CTGGTGAGCG	AGGTGACGGG	CACGCTCACC	GCGGAGGAAC
	6541	TGGTGGCGCG	GCTGGAGGAG	GCGTGCATCG	CCTACGCACG	CCAGCGCACC	GTGCGGGAGT
	6601	TCAGCGAACA	CCCCCAACTG	CGTGACCGTG	GACGCTGGGC	TCCGTTTCGAC	AGCCCGGTCTG
	6661	GTGCGCTGGA	GGGCCTGATC	CCCCCGGTCA	CCTTCCACGG	CGAGCACCCG	CGGCGGCTGG
	6721	GCCGGGTCCC	GGAGCTGGGC	GAGCATAACG	AGTCCGTCCT	GGCGTGCTG	GCCGCGCCCC
40	6781	ACAGCGCCGA	CCGCGAAGAG	GCCGGCCATG	CCGAATGAAC	TCACCGGAGT	CCTGATCCTG
	6841	GCCGCCGTGT	TCCTGCTCGC	CGGCGTACGG	GGGCTGAACA	TGGGCTTGCT	CGCGCTGGTC
	6901	GCCACCTTTC	TGCTCGGGGT	GGTCGCACTC	GACCGAACGC	CGGACGAGGT	GCTGGCGGGT
	6961	TTCCCCGCGA	GCATGTTTCT	GGTGCTGGTC	GCGGTCACGT	TCCTCTTCGG	GATCGCCCGC
	7021	GTCAACGGCA	CGGTGGACTG	GCTGGTACGT	GTCGCGGTGC	GGGCGGTGGG	GGCCCGGGTG
45	7081	GGAGCCGTCC	CCTGGGTGCT	CTTCGGCCTG	GCGGCACTGC	TCTGCGCGAC	AGGCGCGGCC
	7141	TCGCCCCGCG	CGGTGGCGAT	CGTGGCGCCG	ATCAGCGTCG	CGTTTCGCCG	CAGGCACCGC
	7201	ATCGATCCGC	TGTACGCCGG	ACTGATGGCG	GTGAACGGGG	CCGCAGCCGG	CAGTTTCGCC
	7261	CCCTCCGGGA	TCCTGGGCGG	CATCGTCCAC	TCGGCGCTGG	AGAAGAACCA	TCTGCCCCGTC
	7321	AGCGGCGGGC	TGCTCTTCGC	AGGCACCTTC	GCCTTCAACC	TGGCGGTGCG	CGCGGTGTCA
50	7381	TGGTCTCGTC	TCGGGCGGTC	GCGCCTCGAA	CCACATGACC	TGGACGAGGA	CACCGATCCC
	7441	ACGGAAGGGG	ACCCGGGCTT	CCGCCCCGGC	GCGGAACACG	TGATGACGCT	GACCGCGATG
	7501	GCCGCGCTGG	TGCTGGGAAC	CACGGTCTCT	TCCCTGGACA	CCGGCTTCCT	GGCCCTCACC
	7561	TTGGCGGCGT	TGCTGGCGCT	GCTCTTCCCG	CGCACCTCCC	AGCAGGCCAC	CAAGGAGATC
	7621	GCCTGGCCCC	TGGTGCTGCT	GGTATGCGGG	ATCGTGACCT	ACGTCGCCCT	GCTCCAGAGAG
	7681	CTGGGCATCG	TGGACTCCCT	GGGGAAGATG	ATCGCGGCGA	TCGGCACCCC	GCTGCTGGCC

- 31 -

7741	GCCCTGGTGA	TCTGCTACGT	GGGCGGTGTC	GTCTCGGCCT	TCGCCTCGAC	CACCGGGATC
7801	CTCGGTGCCC	TGATGCCGCT	GTCCGAGCCG	TTCCTGAAGT	CCGGTGCCAT	CGGGACGACC
7861	GGCATGGTGA	TGGCCCTGGC	GGCCGCGGCG	ACCGTGGTGG	ACGCGAGTCC	CTTCTCCACC
7921	AATGGTGCTC	TGGTGGTGGC	CAACGCTCCC	GAGCGGCTGC	GGCCCGGCGT	GTACCAGGGG
5	7981	TTGCTGTGGT	GGGGCGCCGG	GGTGTGCGCA	CTGGCTCCCC	CGGCCGCTTG
	8041	GTGGTGCGGT	GAGCGCAGCG	GAGCGGGAAT	CCCCTGGAGC	CCGTTTCCCC
	8101	CTGACGTAGC	GTCAAGTCCA	CGTGCCGGGC	GGGCAGTACG	CCTAGCATGT
	8161	TAATCAGATA	ACCCTGTCCG	ACACGCTGCT	CGCTTACGTA	CGGAAGGTGT
	8221	TGACGAGGTG	CTGAGCCGGC	TGCGCGCGCA	GACGGCCGAG	CTGCCGGGCG
10	8281	GCCGGTGCAG	GCCGAGGAGG	GACAGTTCCT	CGAGTTCCTG	GTGCGGTTGA
	8341	TCAGGTGCTG	GAGATCGGGA	CGTACACCGG	CTACAGCACG	CTCTGCCTGG
	8401	GGCGCCCGGG	GGCCGTGTGG	TGACGTGCGA	TGTCATGCCG	AAGTGGCCCC
	8461	GCGGTACTGG	GAGGAGGCCG	GGGTTGCCGA	CCGGATCGAC	GTCCGGATCG
	8521	GACCGTCTCT	ACCGGGCTGC	TCGACGAGGC	GGGCGCGGGG	CCGGAGTCGT
15	8581	GTTTCATCGAC	GCCGACAAGG	CCGGCTACCC	CGCCTACTAC	GAGGCGGCGC
	8641	ACGCCGCGGC	GGGCTGATCG	TCGTCGACAA	CACGCTGTTC	TTCGGCCGGG
	8701	AGCGGTGCAG	GACCCGGACA	CGGTGCGGGT	ACGCGAACTC	AACGCGGCAC
	8761	CGACCGGGTG	GACCTGGCGA	TGCTGACGAC	GGCCGACGGC	GTCACCCCTG
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	8881	GGCTCCAGAT	GCAGGCGTTC	GACGCCGGCG	GCGGAAGCGC	CCGCCACCTC
	8941	GGGCAGTCGG	AGTCCGCGAA	GCCCCGGAAC	CGGTAGGCGA	TCTCCATCAT
	9001	TCCGTACGCC	GGAAGTCCGC	CACCAGGTGC	GCCCCCGCGC	GGGCGCCCTG
	9061	CAGTTCAGGA	TCGTGCGACC	GGCACCGAAC	GACACGACCC	GGCAGGACGT
25	9121	TTCAGGTGCC	ACGTGACGGG	CTTCTTCTCC	AGCAGGATGA	TGCCGACGGC
	9181	CCGAAGCGGT	CGCCCATGGT	GACGACGAGG	ACCTCATGGG	CGGGATCGGT
	9241	GCAGGTCGGC	GTGCGAGTAG	TGCACGCGCG	TCGCGTTCAT	CTGGCTGGTC
	9301	GTTTCTCGAC	GCGGCTGAGT	TCCTCTCTCC	CCGCGGGTGC	GATCGTCATG
	9361	GCGAGCGCAG	GAAGTCCTCG	TCGGGACCGG	AGTACGCCTC	CCGGGCCCTG
30	9421	AACCCGCCTG	GTACATCAGG	CGGCGCCGAG	GCGAGTCGAC	CGTGGACACC
	9481	ACTCCGGCAG	CGACAGGAGC	GTGGCCGCTT	GCTCGGCCGG	GTAGCACCAG
	9541	GGTGGAACGC	CACCTCGGCA	CGCTCGGGCG	CTGGTTCGTC	GATGAACCGG
	9601	GTGCGAAGTT	CAGCTCCGTG	GCGATCTCGC	GGACGGACTG	CGACTTCGGC
	9661	TGCGGGCCAG	CACGAAGTAC	TCCGCCACAC	CGAGGCGTTC	CAGACGCTCC
35	9721	CGTGGTTCGT	CTTGCTCGCC	ACCGCCTGGA	GGATGCCGCG	GTGCTCGAGC
	9781	CCTCGCGGAT	CTCGTCTGGT	AGGACCACCT	CGTCGTCCTC	CAGCACGGTG
	9841	AGGTGTTGTC	CAGGTCCAGG	ACCAGACACT	TGACAATGGT	CATGGCTGTC
	9901	GGGAGCGCCA	GCGCGTGCTG	GGCCAGCATC	ACCCGGCACA	TCTCGCTGCT
	9961	ATCTCCATGA	GCTTGGCGTC	GCGGTACGCC	CGTTCGACGA	CGTGTCCCTC
40	10021	GCCGACGCGA	GCACCTGTGC	GGCGGTTCGG	GCCCCGGCGG	CGGCTCGTTC
	10081	TGCTTGGCCA	GGATCGTTCG	GGGCACCATC	TCGGGCGAGC	CCTCGTCCCA
	10141	GCGTACTCGC	ACACGCGGGC	CGCGATCTGC	TCCGCGGTCC	ACAGGTCCGC
	10201	GCGACGAGTT	GGTGGTTCGG	GAGCGGCCGG	CCGAACTGCT	CCCGGGTCCG
	10261	ACCGCGGCGG	TGCGGCAGGC	CCGCAGGATC	CCGACGCAGC	CCCAGGCGAC
45	10321	CCGTAGGCGA	GTGACGCCGC	GACCAGCATC	GGCAGTGACG	CGCCGGAGCC
	10381	GCGCCGGCCG	GCACACGCAC	CTGGTCCAGG	TGAGATCGCG	CGTGGCCGGC
	10441	CCGGACGGCT	TCGGGACGCG	CTCGACGCGT	ACGCCGGGGG	TGTCGGCGGG
	10501	ACCGCACGCT	AACCATCCTC	CTGGAGACCG	AAGACGACCA	GGTGGTCCGC
	10561	GACGTGCTCC	AGACCTTGTG	GCCGTGACAG	ACAGCGGTGT	CCCCGTGAGG
	10621	GTCCGCATCG	CCGACAGATC	GCTGCCCCGC	TGCCGCTCAC	TGAAGCCGAC
50	10681	TTCCCGCTGG	TCAGCTCCTT	CAGGAAGGTC	GCCCGCTGAC	CGGCGTCGCC
	10741	ACGGTCCACG	CGGCCATGCC	CTGCGACGTC	ATGACACTGC	GCACGGAATC
	10801	CCGACGTGTG	CGGTGAACTC	GCCGTTCTCC	CGGCTGCCGA	GTCCAGACCC
	10861	GCCGCCACTT	CCGCGCAGAG	CAGGCCGTCG	GCGCCGAGCC	GGACGAGCAG
	10921	AGTTCGCGCG	ACGTGTCCCA	CTCGGCGGCC	CGGTCACCGA	CAAGGTCCGT

- 32 -

"Patent" Sheet

	10981	TCACGCTCAG	GCATCGACGG	CCCGCAGCCG	GTGGACGAGT	GCGACCATGG	ACTCGACGGT
	11041	ACGGAAGTTC	GCGAGCTGGA	GGTCCGGGCC	GGCGATCGTG	ACGTCTGAACG	TCTTCTCCAG
	11101	GTACACGACC	AGTTCCATCG	CGAACAGCGA	CGTGAGGCCG	CCCTCCGCGA	ACAGGTCCGCG
5	11161	GTCCACGGGC	CAGTCCGACC	TGGTCTTCGT	CTTGAGGAAC	GCGACCAACG	CGTGCGCGAC
	11221	GGGGTCGTCC	TTGACGGGTG	CGGTCATGAG	AACACCTTCT	CGTATTTCGTA	GAAGCCCCGG
	11281	CCGGTCTTCC	GGCCGTGGTG	TCCCTCGCGG	ACCTTGCCCA	GCAGCAGGTC	ACAGGGGCGG
	11341	CTGCGCTCGT	CGCCGGTGCG	TTTGTGCAGC	ACCCACAGCG	CGTCGACGAG	GTTGTCGATG
	11401	CCGATCAGGT	CCGCGGTGCG	CAGCGGCCCG	GTCGGATGGC	CGAGGCACCC	CGTCATGAGC
10	11461	GCGTCGACGT	CCTCGACGGA	CGCGGTGCCC	TCCTGCACGA	TCCGCGCCCG	GTCGTTGATC
	11521	ATCGGGTGGA	GCAGCCGGCT	CGTGACGAAG	CCGGGCGCGT	CCCGGACGAC	GATCGGCTTG
	11581	CGCCGACGCG	CCGCGAGCAG	GTCCCCGGCG	GCGGCCATGG	CCTTCTCACC	GGTCCGGGGT
	11641	CCTCGACCGT	CGGGATCAGG	TACGACGGGT	TCATGAAGTG	CGTGCCGAGC	CGACGTGTTT
	11701	AGGTCTTCGG	GCCGGGCCAC	GGAGCTGGCC	AGTTCGTCAA	CCGGGATCGA	CGACGTGTTT
	11761	GTGATGACCG	GGATACCGGG	CGCGCTGCC	GAGACCGTGG	CGAGTACCTC	CGCCTTGACC
15	11821	TCGGCGTCT	CGACGACGGC	CTCGATCACC	GCGGTGGCCG	TACCGATCGC	GGGCAGCGCG
	11881	GACGTGGCCG	TCCGACGAC	ACCGGGGTG	GCCCTGGCGG	GCCCGGCCAC	GAGTTGTGCC
	11941	GTCCGACGTT	CGGTGGCGAT	CCGCGCCCGC	GCCGCGGTAA	GGATCTCCTC	GGACGTGTCG
	12001	ACGAGTGTCA	CCGGGACGCC	GTGGCGCAGC	GCGAGCGTGG	TGATGCCGGT	GGCCATCACT
20	12061	CCCGCGCCGA	GCACGATCAG	CTGGTGGTCC	ACGCTGTTTC	CTCCCTCCGG	GGTCACCATG
	12121	GCAGCGAGTA	CGGGTCGAGG	ACGTCTTCCG	GGGTGACACC	GATCGCGTCC	TTGCGGGCCGA
	12181	GGCCGAGTTC	GTCGGCGAAG	CCGAGCAGCA	CGTCGAACGC	GATGTGGTCG	GCGAACGCGC
	12241	TGCCCCGTGA	GTCGAGGACG	CTCAGGCTGT	CCCGGTGGTC	CGCCGCGGTG	TCCGGTGCCG
	12301	CGCACAGGGC	CGCCAGCGAC	GGGCCGAGCT	CGCGGTCCGG	CAGTTGCTGG	TACTCGCCCT
25	12361	CGGCGCGGGC	CTGCCCCGGA	TGGTCGACGC	AGATGAACGC	GTCGTCGAGC	AGGGTCTTCG
	12421	GCAGTTCGGT	CTTGCCCCGG	TCGTGCGCGC	CGATGGCGTT	CACATGCAGG	TGCGGCAGCC
	12481	GCGGCTCGGC	GGGCAGCACC	GGCCCTTTGC	CCGAGGGCAC	CGAGGTGACG	GTGGACAGGA
	12541	CATCCGCGGC	GGCGGCGGCC	TCCGCGCGAT	CGGTACACCT	GACCGGCAGT	CCGAGGAACG
	12601	CGATGCGGTC	CGCGAACGAC	GCCGCGTGGC	CGGGGTGCGT	GTCGCTGACC	AGGATCCGCT
30	12661	CGATGGGCAG	GACCCTGCTG	AGCGCGTGCG	CCTGGGTCC	CGCCTGTGCG	CCCGCGCCGA
	12721	TCAGCGTGAG	CGTGGCGCTG	TCGGACCGGG	CCAGCAGCCG	GCTCGCGACG	GCGGCGACCG
	12781	GCGCGGTCCG	CATCGCGGTG	ATCACGCCTG	CGTCGCGCAG	GGCGGTGAGA	CTGCCGCTGT
	12841	CGTCGTCGAG	GCGCGACATC	TGCGCGACGA	TCGTGCGCAG	CCGGAAGCGC	GGATAGTTGT
	12901	GCGGACTGTA	CGAAACCGTC	TTCATGCTCA	CGCCGACACC	GGGGACCCGG	TACGGCATGA
35	12961	ACTCGATGAC	GCCGGGAATG	TCCGCGGTCA	GGACGAATCC	GGTACGCGGC	GGCGCCTCGG
	13021	CGAACTCGCC	GCGGCCGAGC	GCGGCGAACC	CGTCGTGACG	CTCGCTGATC	AGCCGGTCCA
	13081	TCATCACGTC	GCGGCCGATC	ACGGAGAGAA	TCCGCTTGAT	GTCACGTTGG	CGCAGGACCC
	13141	TGGTCTGCAT	GTGTCACCTC	CCTTTCTGTT	CCGGAGCTGT	CTTGGTGGTG	CCGCTACGGG
	13201	CGGCTTCCGT	TCTCATCGCA	GCTCCCTGTC	GATGAGGTCG	AAAATCTCGT	CCGCGGTGCG
40	13261	GTCCGCGGAC	AGCACGCCGG	CCGGCGTGGT	CGGGCGGGTC	TCCCGCCGCC	AGCGGTTGAG
	13321	CAGGGCGTCC	AGCCGGGTTT	CGATCGCGTC	CGCCTGGCGG	GCGCCCGGGT	CGACACCGGC
	13381	AACGAGTGCT	TCCAGCCGGT	CGAGCTGCGC	GAGCACCACG	GTCACCGGGT	CGTCCGGGGA
	13441	CAGCAGTTCA	CCGATGCGGT	CGGCGAGTGC	GCGCGGCGAC	GGGTAGTCGA	AGACGAGCGT
	13501	GGCGGACAGT	CGCAGACCGG	TCGCCTCGTT	GAGGCCGTTG	CGCAGCTGCA	CCGCGATGAG
45	13561	CGAGTCCACA	CCGAGTTCCC	GGAACGCCGC	GTCCTCCGGG	ATGTCTCTCC	GGTCCGCGTG
	13621	GCCCAGGACG	GCCGCTGCCT	TCTGCCGGAC	GAGGGCGAGC	AGGTGCGTGG	GGCGTTCTTG
	13681	CTCGTTGCGG	GCGCTCCGGC	GGGCCGACGG	CTTGGGCCGG	CCACGCGACA	GCGGGAGGTC
	13741	CGGCGGCAGG	TCGCCCCGCA	CGGCGACGAC	ACTGCCCCGT	CCGGTGTGGA	CGGCGGCGTC
	13801	GTACATGCGC	ATGCCCTGTT	CGGCGGTGAG	CGCGCTCGCC	CCACCCTTGC	GCATACGGCG
50	13861	CCGGTCCGGC	TCGGTCAGGT	CCGCGGTGAG	GCCACTCGCC	TGGTCCCACA	GCCCCACGCG
	13921	GATCGACAGC	CCTGGCAGCC	CTTGTGCACG	CCGGTGTTCG	GCGAGCGCGT	CGAGGAACGC
	13981	GTTCCGCGCC	GCGTAGTTGC	CCTGACCGGG	GGTGCCCGAG	ACACCGGCCG	CCGACGAGTA
	14041	GACGACGAAT	GCGGCGAGGT	CGGTGTCGCG	GGTGAGCCGG	TGCAGGTGCC	AGGCGGCGTC
	14101	GGCCTTGGGT	TTGAGGACGG	TGTCGATGCG	GTCGGGGGTG	AGGTTGTGCA	GCAGGGCGTC
	14161	GTCGAGGGTT	CCGGCGGTGT	GGAAGACGGC	GGTGAGGGGT	TGAGGGATGT	GGGCGAGGGT

- 33 -

14221 GGTGGCGAGT TGGTGGGGGT CGCCGACGTC GCAGGGGAGG TGGGTGCCGG GGGTGGTGTC
14281 GGGGGGTGGG GTGCGGGAGA GGAGGTAGGT GTGGGGGTGG TTCAGGTGGC GGGCGAGGAT
14341 GCCGGCGAGG GTGCCGGAGC CGCCGGTGAT GACGACGGCC CCCTCGGGGT CCAGCGGCCG
5 14401 CGGGACCGTG AGGACGATCT TGCCGGTGTC CTCGCCGCGG CTCATGGTCG CCAGCGCCTC
14461 GCGGACCTGC CGCATGTCTG GCACCGTCAC CGGCAGCGGG TGCAGCACAC CGCGCGCGAA
14521 CAGGCCGAGC AGCTCCGCGA TGATCTCCTT GAGCCGGTCG GGCCCCGCGT CCATCAGGTC
14581 GAACGGTTCG TGGACGGCGT GCCGGATGTC CGTCTTCCCC ATCTCGATGA ACCGGCCACC
14641 CGGCGCGAGC AGGCCGACGG ACGCGTCGAG GAGTTCACCG GTGAGCGAGT TGAGCACGAC
14701 GTCGACCGGC GGGAACGCGT CGGCGAACGC GGTGCTGCGG GAATCGGCCA GATGCGCTCC
10 14761 GTCCAGGTCC ACCAGATGGC GCTTCGCGGC GCTGGTGGTC GCGTACACCT CCGCGCCAG
14821 GTGCCGCGCG ATCTGCCGGG CGGCGGAACC GACACCGCCG GTGGCCGCGT GGATCAGGAC
14881 CTTCTCGCCG GGGCGCAGCC CGGCGAGGTC GACCAGGCCG TACCAGCGCG TCAGCAACGC
14941 GGTCATCACG GACGCCGCTT GCGGGAACGT CCAGCCGTCC GGCATCCGGC CGAGCATCCG
15001 GTGGTCGGCG ATGACCGTGG GGCCGAAGCC GGTGCCGACG AGGCCGAAGA CGCGGTCGCC
15 15061 CGGTGCCAGA CCGGAGACGT CGGCGCCGGT CTCCAGGACG ATGCCCGCGG CCTCGCCGCC
15121 GAGCACGCCC TGACCGGGGT AGGTGCCGAG CGCGATCAGC ACATCGCGGA AGTTGAGGCC
15181 CGCCGCACGC ACACCGATCC GGACCTCGGC CGGGGCGAGG GGGCGCCGGG GCTCCGCCGA
15241 GTCGGCCGCG GTGAGGCCGT CGAGGGTGCC CGTCCGCGCC GGCCGGATCA GCCACGTGTC
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15361 GCCGCGCAGC CGCAGACGCG GCTCGCCGAG TCGGACGGCG ATGCGCTGCT GCTCGGGGGC
15421 GAGCGTGACG CCGGACTCGG TCTCGACGTG GACGAACCGG CCGGGCTGCT CGGCCTGGGC
15481 GGCGCGCAGC AGTCCGGCCG CCGCGCCGGT GGCGAGGCCG GCGGTGGTGT GCACGAGCAG
15541 ATCCCCGCCG GAGCCGGTCA GGGCGGTGAG CAGCCGGGTG GTGAGCGCAC GCGTCTCGGC
25 15601 CACCGGGTCG TCGCCATCAG CGGCAGGCAA CGTGATGACG TCCACGTCCG TCGCGGGGAC
15661 ATCCGTGGGT GCGGCGACCT CGATCCAGGT GAGACGCATC AGGCCGGTGC CGACGGGTGG
15721 GGACAGCGGG CGGGTGCGGA CCGTCCGGAT CTCGGCGACG AGTTGGCCGG CGGAGTCGGC
15781 GACGCGCAGA CTCAGCTCGT CGCCGTACAG AGTGATCACG GCTCGGAGCA TGGCCGAGCC
15841 CGTGGCGACG AACCAGGGCC CTTTCCAGGC GAACGGCAGA CCCGCGAGCG TGTCGTCCGG
30 15901 CGTGGTGAGG GCGACGGCGT GCAGGGCCGC GTCGAGCAGC GCCGGATGCA CACCGAAACC
15961 GTCCGCCTCG CGGCGCTGCT CGTCCGGCAG CGCCACCTCG GCATACACGG TGTCACCATC
16021 ACGCCAGGCA GCGCGCAACC CTTGGAACGC CGACCCGTAC TCATAACCGG CATCCCGCAG
16081 TTCGTATAG AACCCTGAGA CGTCGACGGC CACGGCCGTG ACCGGCGGGC ACTGCGAGAA
16141 CGGCTCCACA CCGACAACAC CGGGGGTGTC GGGGGTGTCG GGGGTGAGG TGCCGCTGGC
35 16201 GTGCCGGGTC CAGCTGCCCC TGCCCTCGGT ACGCGCGTGG ACGGTCACCG GCCGCCGTCC
16261 GGCTCATCA GCGCTTCCA CGGTACCGA CACATCCACC GCTGCGGTCA CCGGCACCAC
16321 AAGGGGGGAT TCGATGACCA GCTCGTCCAC TATCCCGCAA CCGGTCTCGT CACCGGCCCCG
16381 GATGACCAGC TCCACAAACG CCGTACCGCG CAGCAGGACC GTGCCCCGCA CCGCGTGATC
16441 AGCCAGCCAG GGGTGAGTGC GCAATGAGAT CCGGCCAGTG AGAACAACAC CACCATCGTC
40 16501 GGCGGGCAGC GCTGTGACAG CGGCCAGCAT CGGATGCGCC GCACCCGTCA ACCCGCCGC
16561 CGACAGATCG GTGGCACCGG CCGCCTCCAG CCAGTACCGC CTGTGCTCGA ACGGTACGT
16621 GGGCAGATCC AGCAGCCGTC CCGGCACCGG TTCGACCACC GTGTCCAGT CCACTGCCGT
16681 GCGCAGGGTC CACGCTGCG CCAACGCGT CAGCCACCGC TCCAGCCGC CGTACCGGT
16741 CCGCAACGAC GCCACCGTGT GAGCCTGCTC CATCGCCGGC AGCAGCACCG GATGGGCACT
45 16801 GCACTCCACG AACACCGACC CATCCAGCTC CGCCACCGCC GCGTCCAACG CCACCGGACG
16861 ACGCAGATTC CCGTACAGT ACCCTCATC CACCGGCTCC GTCACCCAGG CGCTGTCCAC
16921 GGTGACACC CACGCCACCG ACGCGGCCCT CCCTGCCACC CCTCCAGTA CCTTGCCAG
16981 TTCATCCTCG ATGGCTTCCA CGTGGGGCGT GTGGGAGGCG TAGTCGACCG CGATACGACG
17041 CACCCGACG CTTTCGGCTT CATAACCGTC CACCACTCC TCCACCGCCG ACGGGTCCCC
50 17101 CGCCACCACC GTCGAAGCCG GGCCGTTACG CGCCGCGATC CACACACCT CGACGAGCC
17161 GACCTACCG GCCGGCAACG CCACCGAAGC CATCGCTCCC CGCCCGGCA GTCGCGCCG
17221 GATGACCTGA CTGCGCAATG CCACCACGCG GGCGGCGTCC TCGAGGTGTA GGGCTCCGGC
17281 CACGACGCG GCCGCGATCT CGCCCTGGGA GTGTCCGATC ACCGCGTCCG GCACGACCCC
17341 ATGCGCCTGC CACAGCGCGG CCAGGCTCAC CGCGACCGCC CAGCTGGCCG GCTGGACCAC
17401 CTCCACCCGC TCCGCCACAT CCGGCCGCGC CAACATCTCC CGCACATCCC AGCCCGTGTG

- 34 -

17461	CGGCAGCAAC	GCCTGAGCGC	ACTCCTCCAT	ACGCGCGGCG	AACACCGCGG	AGTGGGCCAT
17521	GAGTTCCACG	CCCATGCCGA	CCCACTGGGC	GCCCTGGCCG	GGGAAGACGA	ACACCGTACG
17581	CGGCTGGTCC	ACCGCCACAC	CCGTCAACCG	GGCATCGCCC	AGCAGCACCG	CACGGTGACC
17641	GAAGACAGCA	CGCTCCCGCA	CCAACCCCTG	CGCGACCGCG	GCCACATCCA	CACCACCCCC
5	17701	GCGCAGATAC	CCCTCCAGCC	GCTCCACCTG	CCCCCGCAGA	CTCACCTCAC
	17761	CACCGGCAAC	GGCACCAACC	CGTCAACAAC	CGACTCCCCA	CGCGACGGCC
	17821	CTCAAGGATC	ACGTGCGCGT	TCGTACCGCT	CACCCCGAAC	GACGACACAC
	17881	TGCCCAGATC	GACTCGGGCC	ACGGCCTCGC	CTCGGTGAGC	AGCTCCACCG
	17941	CCAGTCCACA	TGCGACGACG	GCTCGTCCAC	ATGCAGCGTC	TTCGGCGCGA
10	18001	CATCGCCATG	ACCATCTTGA	TCACACCGGC	GACACCGCC	GCCGCTGCG
	18061	GTTGCACTTC	AACGAACCCA	GCAGCAGCGG	AACCTCACGC	TCCTGCCCCG
	18121	AATGGCCTGC	GCCTCGATGG	GATCGCCCAG	CGTCGTCCCC	GTCCCGTGGC
	18181	GTCCACATCG	GCGGCGCGCA	GTCCGGCGTT	CACCAACGCC	TGCTGGATGA
	18241	GGACGGGCGG	TTGGGGGCGG	ACAGCCCGTT	GGAGGCACCG	TCCTGGTTCA
15	18301	GCGGACGACC	GCGAGAACGG	TGTGTCCGTT	GCGCTCGGCG	TCGGAGAGCC
	18361	AAGAACGCCG	GCGCCCTCCG	CCCAGCCGGT	GCCGTTGGCG	GCGTCCGCGA
	18421	GCGGCCGTCG	GGGGAGAGTC	CGCCCTGCTG	CTGGAATTCC	ACGAACCCGG
	18481	CATGACGGTG	ACACCGCCGA	CCAGCGCCAG	CGAGCACTCC	CCGTGGCGCA
	18541	GGCCTGGTGC	AGCGCGACCA	GCGACGACGA	GCACGCCGTG	TCCACCGTGA
20	18601	CTGGAGCCCA	TAGAAGTACG	AGATCCGGCC	GGTGAGCACG	CTGGGCTGCA
	18661	GCCGAACCCG	TCCAGGTCCG	CGCCGACGCC	GTACCCGTAC	GAGAAGGCGC
	18721	GCCGGTGTGC	CTGCCGCGCA	GTGTGCCCGG	CACGATGCCC	GCGCTCTCGA
	18781	TGTCGTTTCC	AGCAGGATCC	GCTGCTGGGG	GTCCATGGCC	CGTGCCCTAC
	18841	GCCGAAGAAC	GCGGCATCGA	AGCCGGCGGC	GTCGGAGAGG	AAGCCGCCGC
25	18901	CGATCCGCGG	GTGAGGCCGG	ACGGGTCCCA	GCCACGGTCG	GCCGGGAAGC
	18961	GTGCGCCGCA	CTGTCCACCA	TGCGCCACAG	GTCGTGCGGC	GAGGTGACGC
	19021	TCGGCAGGCC	ATGCCCACGA	TGGCCAGCGG	TTGCTCACGG	GTCGCGGCGG
	19081	AGCGACCGGT	GCGGCACAC	CGAGCAGAGC	CTCGTCCAAC	CGCGACGCGA
	19141	CGTCGGGTAG	TGGAAGACAA	GCGTGGCGGG	CAGTCGGACA	CCGGTCGCCG
30	19201	GTTCCGCGAGT	TCGACGGCGG	TCAGCGAGTC	GATACCCAGT	TCCTTGAAGG
	19261	GGACACGTCC	GCGGCGTCCG	CGTGCCCGAG	CACCGCCGCC	GCGTTGTGCG
	19321	CAGCAGCGCG	GTGTCCCGCT	CAGCGCCGGA	CATGGTGCCG	AGCCGGTCGG
	19381	GGCGGTGGCC	GCCGCCGGGC	GCGATACGGC	GCGGCGCAGA	TCGGCGAAAA
	19441	GTGCGCGGTG	AGGTCCATCG	TGGCCGCCAC	GGCGAACGCG	GTGCCGGTTC
35	19501	TTCCAGCAGG	CGCATGCCCA	CACCGGCCGA	CATGGGGCGG	AAACCGCCGC
	19561	GGTGCGGTTG	GTGCCGCTCA	TGCTGCCGGT	GAGTCCGCTG	TCATCGGCCC
	19621	GGCCAGCGAC	AGCGCGGGCA	GTCCTTCGGC	ATGGCGCAGC	GTCGCGAGTC
	19681	CCCGTTCCGC	GCCGAGTAGT	TGCCCTGGCC	GCGGCCGCCC	ATGATGCCCG
	19741	GTAGAGGACG	AACGAGCGCA	GGTCCGCGTC	CCGGGTCAGC	TCGTGCAGGT
40	19801	GTCGGCTTTG	GGGCGCAGTG	TGGTGGCGAG	CCGCTCCGGG	GTGAGTGCCG
	19861	GTCGTGAGC	ACGGCTGCCG	TGTGGAAGAC	CGCCGTGAGC	GGCCTGCCCG
	19921	CGCGGCGGCG	AGCTGGTCCC	GGTGGCGGAC	GTCACAGCGG	ATGTGGACAC
	19981	CGCCGGCGGT	TCGTGCGCG	ACAGCAACAG	GAGGTGGCGG	GCGCCATGCT
45	20041	ATGCCGGGCG	AGGAGACCTG	CCAGCACACC	CGAGCCGCGG	GTGATGACCA
	20101	CGGGTCGAGC	AGCGGTTCCG	GCGTTTCCCG	GGCGGCCGTG	CGGGTGAACC
	20161	GTACCGGGCG	TCGGTGACGC	GGACGTACCG	CTCGGCCAGT	GTCGTGGCGG
	20221	CTCGATGGGG	GTGTGCGTGC	CGGTCTCCAC	CAGCACGAAC	CGGCCCGGGT
	20281	GGCGGACCGG	ACGAGGCCGG	CGACCGCTCC	TCCGACCGGT	CCCGCGTCGA
	20341	GAGGGTGGTC	TCCGCAGGGC	CGTCCTCGGC	GATCACCCGG	TGCAGCTCGC
50	20401	CTCGGTGAGC	CGGTACGTCT	CGTCGAGGAC	ATCCGCGCCC	GGTTCCGGGA
	20461	GATGTGGACC	GCGTCCGCG	GACCGGGCCC	GGGAGTGGGC	AGCTCGGTCC
	20521	GTACAAGGAG	TTCCGTACGA	CGGCGGCGTC	GCCGTCGACG	TTCACCGGTC
	20581	CGCGGCGACG	GTCACCACCG	GTTGGCCGAC	CGGGTCCGTC	GCATGCACGG
	20641	CGGGCCCTGA	GTGATCGTGA	CGCGCAGCGT	GGTGGCCCCG	GTCGTGTGGA

- 35 -

20701 GCTCCACGAG AACGGCAGCC GCACCTCCGC TTCCTGTTCC GCGAGCAGCG GCAGGCAGGT
20761 GACGTGCAAG GCCGCGTCGA ACAGCGCCGG GTGGACGCCA TAGTGCGGCG TGTCGTCCGC
20821 CTGTTCCCGG GCGATCTCCA CCTCGGCGTA CAGGGTTTCG CCGTCGCGCC AGGCGGTGCG
20881 CAGTCCCTGG AACGCTGGGC CGTAGCTGTA GCCGGTCTCG GCCAGCCGCT CGTAGAACGC
5 20941 GCTCACGTCG ACGGTCGCG CGCCCGGCGG CGGCCACGCG GCGGCGGGA CCGCCGCGAC
21001 GCTTCCGGCC CGGCCGAGGG TGCCGCTGGC GTGCCGGGTC CAGCTGTCCG TGCCCTCGGT
21061 ACGCGCGTGG ACGGTCACTC GCCGCCGTCC GGCTCATCG GCCCCTTCGA CCGTCACCGA
21121 CACATCCACC GCGCCGGTCA CCGGCACCAC GAGCGGGGTC TCGATGACCA GTTCATCCAC
21181 CACCCCGCAA CCGGTCTCGT CACCGGCCCG GATGACCAGC TCCACAAACG CCGTACCCGG
10 21241 CAGCAGAACC GTGCCCGCA CCGCGTGATC AGCCAGCCAG GGATGCGTAC GCAACGAGAT
21301 CCGGCCAGTG AGAACACAC CACCACGTC GTCGGCGGGC AGTGCTGTGA CCGCGGCCAG
21361 CATCGGATGC GCCGCCCGG TCAACCGGTA GGTGGGACAG TCGAGCAGCC GTCCCGGCAC
21421 CAGCCAGTAC CGCCTGTGCT CGAACCGGTA GGTGGGACAG TCGAGCAGCC GTCCCGGCAC
21481 CGGTTCGACC ACCGTGTCCC AGTCCACTGC CGTGCCGAGG GTCCACGCCT GCGCCACGCG
15 21541 CGTCAGCCAC CGTCCCAGC CGCCGTACC GGTCCGCAAC GACGCCACCG TGTGAGCCTG
21601 TTCCATCGCC GGCAGCAGCA CCGGATGGGC GCTGCACTCC ACGAACACGG ACCCGTCCAG
21661 CTCCGCCACC GCCGCGTCCA GCGCAGCGGG GCGACGAGG TTCCGGTACC AGTAGCCCTC
21721 ATCCACCGGC TCGGTCACCC AGGCGCTGTC CACCGTGGAC CACCAGGCCA CCGACCCGGT
21781 CCCGCCGAA ATCCCTCCA GTACCTCGGC CAACTCGTCC TCGATGGCTT CCACGTGGGG
20 21841 CGTGTGGGAG GCGTAGTGA CCGCGATACG GCGCACTCGC ACGCCTTCGG CCTCGTACCG
21901 CGTCACCACT TCTTCCACCG CGGACGGGTC CCCCGCCACC ACAGTCGAAG ACGGGCCGTT
21961 ACGCGCCGCG ATCCACACGC CCTCGACCAG GTCCACCTCA CCGGCCGGA ACGCCACCGA
22021 AGCCATCGCC CCCCGCCCGG CCAGCCGCCG GCGATCACC TGGCTGCGCA AGGCCACCAC
22081 GCGGGCGGCG TCCTCAAGGC TGAGGGCTCC GGCCACACAC GCCGCCGCGA TCTCGCCCTG
25 22141 GGAGTGTCG ACCACCGCGT CCGGCACGAC CCCATGCGCC TGCCACAGCG CCGCCAGGCT
22201 CACCGCGACC GCCCAGCTGG CCGGCTGGAC CACCTCCACC CGTCCGCCA CATCCGGCCG
22261 CGCCAACAT TCCCGCACAT CCCAGCCCGT GTGCGGCAAC AACGCCCGCG CACACTCCTC
22321 CATAACGACC GCGAACACCG CAGAACACGC CATCAACTCC ACACCCATGC CCACCCACTG
22381 AGCACCCCTGC CCGGAAAGA CGAACACCGT ACGCGGCTGA TCCACCGCCA CACCCATCAC
30 22441 CCGGGCATCG CCCAACAAACA CCGACGGTG ACCGAAGACA GCACGCTCAC GCACCAACCC
22501 CTGCGCGACC GCGGCCACAT CCACACCACC CCGCGCAGA TACCCCTCCA GCCCTCCAC
22561 CTGCCCCCGC AGACTCACCT CACTCGAGC CGACACCGGC AACGGCACCA ACCCATCGAC
22621 AGCCGACTCC CCACGCGACG GCCCGGGAAC ACCCTCAAGG ATCACGTGCG GTTTCGTACC
22681 GCTACCCCG AAAGCGGAGA CACCGGCCCG GCGCGGACGT CCGCGCTCGG GCCACGCCCG
35 22741 CGCCTCGGTG AGCAGTTCCA CCGCGCCCTC GGTCCAGTCC ACATGCGACG ACGGCTCGTC
22801 CACATGCAGC GTCTTCGGCG CGATGCCATA CCGCATCGCC ATGACCATCT TGATGACACC
22861 GCGACACCC GCAGCCGCT GCGCATGACC GATGTTGAC TTCAACGAAC CCAGCAGCAG
22921 CGGAACCTCA CGCTCCTGCC CGTACGTCG CAGAATCGCG TGCGCCTCGA TGGGATCGCC
22981 CAGCGTCGTC CCGTCCCGT GCGCCTCCAC CACGTCCACG TCGGCGGGGG CGAGCCCCGC
40 23041 CTTGTGGAGG GCCTGGCGGA TGACGCGCTG CTGGGAGGGG CCGTTGGGTG CCGAGATGCC
23101 GTTGGAGGCG CCGTCCCTGGT TGACGGCGGA GGAGCGGACG ACCGCGAGGA CCGTGTGTCC
23161 GTTGCGCTCG GCGTCGGAGA GCTTTTCGAC GACGAGGACG CCGGCCCCCT CGGCGAAACC
23221 GGTGCCGTCC GCCGCGTCAG CGAACGCCTT GCACCGTCCG TCCGGCGCGA CGCCGCCCTG
23281 CCGGGAGAAC TCCACGAAGG TCTGTGGTGA TGCCATCACT GTGACACCAC CGACCAGCGC
45 23341 CAGCAGAC TCCCCGGTCC GCAGCGCCTG CCGGCCCTGG TGCAGCGCGA CCAGCGACGA
23401 CGAACACGCC GTGTCGACCG TGACCGCCG ACCCTCCATG CCGAAGAAGT ACGACAGCCG
23461 TCCGGCGAGC ACCGCGGGCT GTGTGCTGTA GGCGCCGAAT CCGCCAGGT CCGCGCCCGT
23521 GCCGTAGCCG TAGTAGAAGC CGCCGACGAA GACGCCGGTG TCGCTGCCG GCAGGGTGTC
23581 CGGCACGATG CCGGCGGTGTT CGAGCGCCTC CCAGGCGATT TCGAGGAGGA TCCGCTGCTG
50 23641 CGGGTCGAGT GCGGTGGCCT CGCGCGGACT GATGCCGAAG AACCGGCGAT CGAAGTCGGC
23701 GCGCGCCGCG AGTGCGCCGG CCCGCCCGGT GGCGGACTCG GCGGCGGCGT CGAGCGGGC
23761 CACGTCCCAG CCGCGGTGCG TGGGGAAGTC GCCGATCGCG TCGCGGCCGT CCGCGACGAG
23821 CTGCCACAGC TCTTCCGGTG AGGTGACGCC GCCCGGCAGT CCGCAGGCCA TGCCGACGAC
23881 GCGGAGCGGC TCGTTCGCCG CGGCGCGCAG CGCGGTGTTC TCCCGGCGGA GCTGCGCGTT

- 36 -

23941	GTCCTTGACC	GACGTCCGCA	GCGCCTCGAT	CAGGTGCTTC	TCGGCCATCG	CCTCATCCCT
24001	TCAGCACGTG	CGCGATGAGC	GCGTCTGCGT	CCATGTCGTC	GAACAGTTCG	TCGTCCGGCT
24061	CCGCGGTCTG	GGTGCTCGCG	GGTGCTGTG	CCGGTGGTTC	ACCGCCGTCC	GGGGTCCCGT
24121	TGTCGTCCGG	GGTCCCGTTG	ACGTCCGGGG	CCAGGAGGGT	CAGCAGATGA	CGGGTGAGCG
5	24181	CGCCGGCGGC	GGGATAGTCG	AAGACGAGCG	TGGCCGGCAG	CGGAATGCCG
	24241	AGAGCCGGTT	GCGCAGGCCG	AGCGCGGTGA	GCGAGTCGAC	CCCGAGGTCC
	24301	TGGTGGCCGT	GACCGCCGCC	GCGTCGGTGT	GGCCCAGCAG	GGTGGCGGCG
	24361	CGACGCCGAG	CAGCACCTGT	TCCCGTTTCT	TGTGGGGCAG	GTCCGGCAGG
	24421	GGGAGCCGCC	GTCGGTCGCG	GAGCGCCGGG	TGGGGCGCTG	GATCGGTTCG
10	24481	ACGGGTTCGC	GGGCCCGGGT	GGGGCGGTTC	CCACGACCAC	GGCTTCCCCG
	24541	CGCGCTCGAG	GAGGTCGGTC	AGCCGGTCCG	CCGCGGCGGT	GAACGCCACG
	24601	CTTGTCGCGG	GCGCAGGTTC	GCCAGGGCCT	GGAGCGGTTC	GGCCGCCTCG
	24661	CGGCGAGAAC	GAACGCGGTC	AGGTTCGAGT	CGCGGGTCAG	GCGGTGCAGT
	24721	ACTCGGCGGT	GCCGTCCGCG	TGGACGACCG	CGGTACCCGG	GGTTTCCGGC
15	24781	GCTCGTACCG	GATCACTTCG	GCGCCGTGTC	CGCCGAGGTG	TCCGGCGAGT
	24841	CGCCCGCGAG	GAGGACGGTG	TCGCCGTACG	AGGCCGCGGC	CGTGGTGGGC
	24901	CGAGGCGGGG	CGCTTCGAGG	CGCCCGTCGG	CCAGGCGCAG	GTGCGGTTTC
	24961	AGAGGGCGGC	GGCGCGCGCG	GGGGTGACCG	TGTCGGTGGT	CTCCACGAGC
20	25021	CCGGTTCCGC	GGTGTTCGAG	AGTGCGGCGA	CGGCACCGGC	GACGGGCCCC
	25081	ACACCACCAG	CGTGGCGCCG	GCGGTCCTCG	GGTCGTCCAG	TGCGGTACCG
	25141	GACCGGATAC	CGGGACGACG	ATGACGTCGG	GCGTGGCGTC	GTCGCCGAGG
	25201	GGCGGGCCGT	GGTGCCGGGT	GCCGCCGGGG	CCCGGACGCC	GGTCCAGGTG
	25261	GCCGCACGTC	CCCGTCCGGG	CCCGTCCGTG	CGGGGGGGCC	GGTGATGAGC
25	25321	GAGCCACCGG	CCGTCCCAGT	TCGTCCGGCG	GGTGCACGCG	GGCGCCGCCC
	25381	CGTGGACGAA	GGTGACGCGC	AGTTTTCGTG	CGCCGCTGGT	GTGGACACCG
	25441	ACGCGAACCG	CAACCGTACC	CCCGCGTTCT	CGGCGGCCGC	GCCGATGCTG
	25501	GCGCGGTGAC	GAGCAGCGCC	GGGTGCAGTG	TGTAGCGGGC	GGCGTCCCTG
	25561	CGTCGAGGGC	GACTTCGGCG	CAGACGGTGT	CTCCGTGGCT	CCACGCGGCG
30	25621	GGAACTCCGG	GCCGAACCTC	TATCCCGCGT	CGTCGAGTCG	CTGGTAGAAG
	25681	CGACCGGTTT	CGCGTGCTCG	GGCGGCCAGG	GCCCCGGCGT	GGTGGCCGGT
	25741	CGATGCCGGC	GAAGCCGGAG	GCGTGGCGGG	TCCATGTCCG	GTCGCCGTCC
	25801	GGACGCGCAC	GGCAGCGCGT	CCCGTGTGTC	CGGGCGCGGC	GACGGTCACG
	25861	CGGCGCCGGT	GGCGGGCAGG	ACGAGCGGTG	TCTCGACGAC	CAGTTCGTTC
	25921	AGCCTGCCTC	GTCGGCGCCG	CGTCCGGCCA	ATTCCAGGAA	GGCGGGTCCG
35	25981	CGGCGCCGTC	GACGGAGTGA	CCGGCCAGCC	ATGGGTGGGT	GGCCAGCGAG
	26041	TGAGCAGCAC	CTCGTCGGAG	TCGGGGAGCG	CCACCGACGC	GGCGAGCAGC
	26101	CGGCGTCGAG	TCCGAGGCCG	GAAGCGTCCG	TGCCGGCCGC	GGTCTCGATC
	26161	CATGGTGGAA	GGCGTATGTG	GGCAGGTTCG	GTGCCGTTCG	CGTCGCGGGG
	26221	CCCAGTCGAC	GGGCACGCCG	GTTGTGTGCG	CCTCGGCCAG	CGCGGTGAGC
40	26281	CTCCCCCGCC	GCGGCGGAGC	GTGGCGACGG	TCGCGCCGTC	GATCGCGGGC
	26341	GGTGC GCGCT	GACCTCGACG	AACACGGTGT	CACCCGGCTC	GCGGGCAGCG
	26401	TGGCGAAGCC	TACGGGGTGG	CGCATGTTGC	GGAACAGTA	CTCGTCGTTC
	26461	CGATCCAGCG	TTCGTTCGGC	GTGGAGAACC	ACGGGATCTC	GGGCGTTCGC
	26521	CCGCGACGAT	CCGCTGGAGT	TCGTCTGACA	GCGGGTCGAC	GAACGGGGTG
45	26581	AGTCGACGGC	GATGCGGCGC	ACCCAGACGC	CGCGGGCCTC	GTAGTCGGCG
	26641	CGACGGCGTC	CGGGCGCCCC	GCGACGGTCG	TGGTGGTGGC	GCCGTTGCGG
	26701	AGACGCCGTC	GATCCGGGCG	GCATCCGCCT	CGACGTCGGC	GGCCGGGAGC
	26761	CCATCGCGCC	GCGTCCGGCG	AGTTCCGCGA	GGAGCAGGAG	AACGCTGCGC
	26821	GGCGGGCACC	GTCCTCCAGG	GTGAGCGCTC	CGGCGACACA	GGCCGCGGCG
50	26881	GGGAGTGTCC	GATGACGGCG	TCCGGGCGTA	CGCCCGCGGC	CTCCACACAG
	26941	ACACCATGAC	GGCCCAGCAG	ACGGGGTGCA	CGACGTCGAC	GCGGCGGGTC
	27001	CGTCGAGCAT	GGCGATGGGG	TCCCAGCCCC	TGTGCGGGAT	CAGCGCGTCG
	27061	GCATCCTGGC	GGCGAACACC	GGGGAGGCGC	CCATCAGTTC	GACGCCCATG
	27121	GCGGTCCTTG	TCCGGGGAAG	ACGAAGACGG	TGCGCGGCTC	GGTGAGCGCC

- 37 -

27181 CGACGTCGTC GTCGAGCAGC ACGGCGCGGT GCGGGAACGT CGTACGCCTG GCGAGCAGGC
27241 CCGCGGCGAT GGCGCGCGGG TCGTGGCCGG GACGGGCGGC GAGGTGCTCG CGGAGTCGGC
27301 GGACCTGGCC GTCGAGGGCC GTGGCGGTCC GCGCCGAGAC GGGCAGTGGT GTGAGCGGCG
27361 TGGCGATCAG CGGCTCACCG GGCTTCGAGG CCGACGGCTC CTCGGCCGGC GGCTCCCCCG
5 27421 CCGGGTGGGC TTCCAGCAGG ACGTGGGCGT TGGTGCCGCT GACGCCGAAG GAGGACACAC
27481 CGGCGCGCCG CGGGCGGTCT GTCTCGGGCC AGGGCCGGGC ATCGGTGAGG AGTTCGACGG
27541 CGCCGGCCGT CCAGTCGACG TGCAGGAGCG GCGTGTCCAC GTGCAGGGTG CGCGGCAGGG
27601 TGCCGTGCCG CATGGCGAGG ACCATCTTGA TGACACCGGC GACACCCGCG GCGGCCTGAG
27661 TGTGGCCGAT GTTGACTTC AGCGAGCCCA GCAGACCGG GGTGTGCGCG CCCTGCCCCG
10 27721 AGGTGGCCAG CACCGCCTGT GCCTCGATGG GATCGCCCAG CCTGGTGCCG GTGCCGTGCG
27781 CCTCCACGGC GTCCACGTTC GCCGGGGTGA GCCCGGCGTT GGCCAGGGCC TGCCGGATCA
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27961 TCTCGACGAT CAGCACACCG GACCCCTCGG CGAAACCGGT GCCGTGAGCC GCATCCGCGA
15 28021 ACGCCTTGCA GCGCGCGTCG GGCGCGAGAC CCCGCTGCTG GGAGAACTCG ACGAAGCCGG
28081 ACGGCGAGGC CATCACCGTG ACGCCGCCGA CCAGGGCGAG CGAGCATTCG CCGGAGCGCA
28141 GTGACTGCCC GGCTTGGTGC AGCGCCACCA GCGACGACGA ACACGCCGTG TCGACCGTGA
28201 CCGCCGACC CTCCAGACCG TAGAAGTACG ACAGCCGACC GGACAGCACA CTGGTCTGGG
28261 TGCCGGTTCG GCCGAAACCG CCCAGGTCGG TGCCGAGTCC GTACCCGTGCG GAGAAGGCGC
20 28321 CCATGAACAC GCCGGTGTGCT CTCGCGCA GCGACTCCGG GAGGATCCCG GCGTGTTCGA
28381 GCGCCTCCCA CGAGGTCTCC AGGACCAGAC GCTGCTGCGG GTCCATCGCC AGCGCCTCAC
28441 GCGGACTGAT CCCGAAGAAC GCCGCGTCGA AGTCCGCCAC CCCGGCGAGG AAGCCACCAT
28501 GACGCACGGT CGACGTGCCC GGATGATCCG GATCGGGATC GTACAGCCCG TCCACGTCCC
28561 AACCACGGTC CGTCGGAAC GCGGTGATCC CGTCACCACC CGACTCCAGC AGCCGCCACA
25 28621 AGTCCTCCGG CGACGCGACC CCACCCGGCA GCCGGCAGGC CATCCCCACG ATCGCCAACG
28681 GTCGTCCTG CCGGACGGCC GCGGTCTGGG TGCGGGTCGG CGATGCCGTC CGGCCGGACA
28741 GCGCCGCGGT GAGCTTCGCC GCGACGGCGC GCGGCGTCGG GAAGTCGAAG ACCGCGGTGG
28801 CGGGCAGCCG TACGCCCCGT GCCTCGGTGA AGGCGTTGCG CAGCCGGATC GCCATGAGCG
28861 AGTCGACGCC GAGTTCCTTG AACGTGGCGG TCGCCTCGAC CCGTGCGGCA CCGTCGTGGC
30 28921 CGAGTACGGC CGCGGTGCAC TGCCGGACGA CCGCGAGCAC GTCCTTTTCG GCGTCCGCGG
28981 CGGAGAGCCG CGCGATCCGG TCGGCGAGGG TGGTGGCGCC GGCCGCCCCG CGCCGCGGCT
29041 CCCGGCGCGG TGCGCGCAGC AGGGGCGAGC TGCCGAGGCC GGCCGGGTGCG CCGGAGACCA
29101 GCGCCGGGTC CGAGGACCGC AACGCCGCGT CGAACAGCGT CAGTCCGCCT TCGGCGGTCA
29161 GCGCCGTCAC GCCGTGCGGG CGCATGCGGG CGCCGGTGCC GACCGTCAGC CCGCTCTCCG
35 29221 GTTCCACAG GCCCCAGGCC ACGGACAACG CCGGAGTCC GGCTGCCCGG CGTGTTCGG
29281 CCAGCGCGTC GAGGAACGCG TTCGCGGCCG CGTAGTTGCC CTGTCCGGGG CTGCCGAGCA
29341 CACCGGCGGC CGACGAGTAG AGGACGAACG CCGCCAGTTC CGTGTCTGCG GTGAGTTCGT
29401 GCAGGTGCCA CGCGGCGTCC ACCTTCGGGC GCAGACCGT CTCGAGCCGG TCGGGGGTGA
29461 GCGCGGTGAG GACGCCGTCG TCGAGGACGG CCGCGGTGTG CACGACGGCC GTGAGCGGGT
40 29521 GCGCCGGGTC GATCCCCGCC AGTACGGAGG CGAGTTCGTC CCGGTCGGCG ACGTCGCAGG
29581 CGATCGCCGT GACCTCGGCG CCGGGCACGT CGCTCGCCGT GCCGCTGCGC GACAGCATCA
29641 GCAGCCGGCG CACGCCGTGG CGTTCGACGA GGTGGCGGCT GATGATGCCG GCCAGCGTCC
29701 CGGAGCCACC GGTGACGAGC ACGGTGCCGT CCGGGTCGAG CGCCGAGCG TCACCCGCCG
29761 GGACCGCCG GGCCAGACGG CCGGCGTACA CCTGGCCGTC ACGCAGCACC ACCTGGGGCT
45 29821 CATCGAGCG GGTGGCCGCT GCGAGCAGCG GCTCGGCGGT GTCCGGGGCG GCGTCGACGA
29881 GGACGATCCG GCCGGGGTGT TCGGCCTGCG CGGTCCGCAC CAGTCCGGCG GCCGCGGCCG
29941 ACGCGAGACC GGGCCCGGTG TGGACGGCCA GGACCGCGTC GGCGTACCGG TCGTCGGTGA
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50 30121 GGCCGGTCTG CGCGGTCTG GCGGCGAGCT CCGGGAGCTC GGCCAGCACC GGGCGACGA
30181 GGCCCGGAAC GGCTCCCGTG ATCGTCAGGG GCGCCTGCG CACGCGCCG ATGGTGGCGA
30241 CCGGCCCCGCC GGTCTCGTCC GCGAGGTGTA CGCCGTCAGC GGTGACGGCG ACGCGTACCG
30301 CCGTGGCGCC GGTGGCGTGG ACGCGGACGT CGTCGAACGC GTACGGAAGG TGGTCCCCCTT
30361 CCGCGGCGAG GCGGAGTGCG GCGCCGAGCA GCGCCGGGTG CAGGCCGTAC CGTCCGGCGT

- 38 -

30421	CGGCGAGCTG	TCCGTGCGCG	AGGGCCACTT	CCGCCCAGAC	GGCGTCGTG	TCGGCCCAGA
30481	CGGCGCGCGG	GCGGGGCAGC	GCGGGCCCGT	CCGTGTACCC	GGCTCGGGCC	AGACGGTCCG
30541	CGATGTCGTC	GGGGTCCACC	GGCCGGGCGG	TGGCGGGCGG	CCACGTCGAC	GGCATCTCCC
5	30601	GCACGGCCCG	GGCCGTCCGC	GGGTGCGGGG	CGAGGATTCC	GTGCGCGTGC
	30661	CCCCCGCCGC	GTGCCGCGTG	TGCACGGTGA	CCGCGCGGCG	GCCGTCCGCC
	30721	TCACCGTGAC	GGAGAGCGCG	AGCGCACCGG	ACCGCGGCAG	CGTGAGGGGG
	30781	TGAACGTGTC	GAGGGCGCCG	CAGCCGGCTT	CGTCGCCCCG	CCGGATCGCC
	30841	GGGCCGCGGC	GGGCAGCACC	GCGAGGCCGT	GCAGGGAGTG	CGCCAGCGGA
	30901	CGACCCGGCC	GGTGAGCACC	AGGTGCGCGG	TGCCGGGCAG	GGTGACCGCC
10	30961	CCGGGTGCGC	GACCGGCGTC	TGTCCGGCCG	GGGCCGCGTC	GCCCCGCGCA
	31021	GCCAGTAGCG	GACCCGCTCG	AACGGGTACG	TCGGCGGGTG	CGAGGCGCGT
	31081	GGTCGATGAC	CTTCGGCCAG	TCGACCGTGA	CGCCGTGCGT	GTGCAGCCGG
	31141	TCAGGGCGGA	TCGCGGTTTC	TCGTGCGCGT	GCAGCATCGG	GATGCCGTGC
	31201	TCAGGCTCCG	GTCCGGGCGG	ATCTCCAGGA	GCACCGCCCC	GTCGTGCGCG
15	31261	CCCCGAACCG	GACGGTGTGC	CGGACCTGTC	GTACCCAGTA	CTCCGGCGTG
	31321	CGCCGCGGCG	CATCGGGATC	CTCGGCTCGT	GGTACGTCAG	GCTCTCCGCG
	31381	ACTCCTCGAG	CATCGGCTCC	ATCCGCGCCG	AGTGGAACGC	GTGGCTGGTC
	31441	TGAAGCGGCC	GAGCCGGGCC	GCGACGTCGA	GCACCGCCTC	CTCGTCACCG
20	31501	TCGACGCGGG	CCCGTTGACC	GCGGCGATCT	CCACGCCGTC	CCGCAGCAGC
	31561	CCCGTTCCGA	CGCGATCACG	GCGGCCATCG	CCCCGCCGGA	CGGCAGCGCC
	31621	GGGCCCGTGC	GGACACCAGC	CTGCACGCGT	CCTCCAGGGA	CCAGACGCCG
	31681	CGGCGGCCAG	CTCGCCGATC	GAATGGCCCA	CGAAGGCGTC	CGGGCGTACG
	31741	CGAGCTGTGC	GCCGAGTGCG	ACCTGGAGCG	CGAACACCGC	GGGCTGGGCG
25	31801	CGTGAGGTC	GAGCCCGGCG	GGCACGTCGA	GGGCGTCCAG	CACCTCGCGG
	31861	CGAAGACGTC	GTAGGCGGCG	GCCAGTCCGT	CGCCCATGCC	GGGACGTTGT
	31921	CGGAGAAGAG	CCACACGAGG	CGGCGGTCCG	GTTCTGCGGC	GCCGGTGACC
	31981	CGATCAGCGC	GGCCCGGTGC	GGGAAGGCCG	TGCGGGCGAG	CAGGGCCGCG
	32041	GCTCGTCCTC	CTCGCCGGTG	GCGAGGTGGG	CGCGCAGGCG	GTGTACCTGT
30	32101	CCTGCGGGGT	GCGTGCCGAG	AGCAGCAGGG	CGAGCGGTCC	GGTGTGCGGT
	32161	GTTGCGGGGC	CGGTGCGGGG	TGGCTTTTCGA	GGATGATGTG	AGCGTTGGTG
	32221	CGAAGGAGGA	CACCCCGGCG	CGCCGTGGGC	GGTCGGTTTC	GGGCCAGGGG
	32281	TGAGGAGTTC	GACGGCGCCG	GCCGTCCAGT	CGACGTGCGA	GGACGGCGTG
	32341	GGGTGCGCGG	CAGGGTGCCG	TGCCGCATGG	CGAGGACCAT	CTTGATGACA
35	32401	CCGCGGCGGC	CTGAGTGTGG	CCGATGTTGG	ACTTCAGCGA	GCCCAGCAGC
	32461	CGCGATGCTG	CCCGTAGGTG	GCCAGTACCG	CCTGCGCCTC	GATGGGGTCG
	32521	TCCCGGTGCC	ATGCGCCTCG	ACAGCGTCCA	CATCCGCCGG	GGTGAGCCCG
	32581	GCGCCTGCCG	GATCACCCGC	TCCTGCGACG	GCCCGTTCGG	CGCCGACAAC
	32641	CACCGTCCTG	GTTGACCGCC	GAACCACGCA	CGACCGCCAG	GACATTGTGG
40	32701	CGGCGTCGGA	GAGCCTCTCG	ACGATCAGCA	CACCGGATCC	CTCGGCGAAA
	32761	CAGCCGCATC	CGCGAACGCC	TTGCAGCGGC	CGTCCGGGGA	GAGGCCCCGC
	32821	AGTCCACGAA	GCCGGACGGC	GAGGCCATCA	CCGTGACGCC	GCCGACCACG
	32881	ACTCCCCCGA	GCGCAGCGAC	TGCCCCGGCT	GGTGCAGCGC	CACCAGCGAC
	32941	CGTGTTCCAC	CGTGACCGCC	GGACCCTCCA	AACCGTAGAA	GTACGACAGC
45	33001	GCACACTGGT	CTGGGTGCTG	GTGGCACCAG	AACCGCCGCG	GTCGGCTCCA
	33061	CGTAGAAGTA	GCCGCCCATG	AACACGCCGG	TGTCGCTTCC	GCGCAGCGAC
	33121	TCCCGGCGTG	TTCCAGCGCC	TCCACAGGAG	TCTCCAGGAC	CAGACGCTGC
	33181	TCGCCAGCGC	CTCACGCGGA	CTGATCAGGA	AGAACGCCCG	GTCGAAGTCC
	33241	CGAGGAAGCC	ACCATGACGC	ACGGTGCAGC	TGCCCCGATG	ATCCGGATCC
	33301	GCCCCGTCCAC	GTCCCAACCA	CGGTCCGTCG	GAAACGCCGT	GATCCCGTCA
50	33361	CCAGCAGCCG	CCACAAGTCC	TCCGGCGACG	CGACCCACC	CGGCAGCCGG
	33421	CCACGATCGC	CAACGGCTCG	TCCTGCCGGA	CGGCCGCGGT	CGGGGTACGC
	33481	TGGCCCGCGC	GCCGGCCAGT	TCGTCCAGGT	GGGCGGCGAG	CGCCTGCGCC
	33541	CGAAGACGAG	CGTAGCGGGC	AGCGTCAGGC	CCGTGCGGTC	GGCCAGCCGG
	33601	CGACGCCGGT	CAGCGAGTCG	AAGCCCACTT	CCCTGAACGC	GCGCGCGGGT

- 39 -

5 33661 GGGCGTCCGG GTGGCCGAGC ACCGCGGCAG CGCTGGTACG GACGAGGTCG AGCATGTCCG
33721 GCGCGGCCCG AGGTGCGGAC GTGCGCCGGA CGGCCGGCAC GAGGGTGCGT AGGACCGGCG
33781 GGACCCGGTC GGACGCGGCG ACGGCGGCGA GGTGAGCCG GATCGGCACG AGCGCGGGCC
33841 GGTGCGGTGTG CAGGGCCGCG TCGAACAGGG CGAGCCCCTG TCGGGCCGTC ATCGGGGTCA
33901 TGCCGTTGCG GGCGATGCGG GCCAGGTCGG TGGCGGTGAG CCGCCCGCCC ATCCCGTCCG
33961 CCGCGTCCCA CAGTCCCCAG GCGAGCGAGA CGGCGGGCAG CCCCTGGTGG TGCCGGTGCG
34021 GGGCGAGCGC GTCGAGGAAC GCGTTGCCGG TCGCGTAGTT GGCGTGACCC GCGCCGCCGA
34081 ACGTGCGGGA TATGGACGAG TACAGGACGA ACGCGGCCAG GTCGAGATCG CGCGTCAGCT
10 34141 CGTGCGAGTG CCAGGCGACG TCCGCTTGA CCCGCGACAC GCGGTCCCAC TGCTCCGGCC
34201 GCATGGTCGT CACGGCCGCG TCGTCGACGA TCCCGGCCAT GTGCACGACG GCGCGCAGCC
34261 GCTGGGCGAC GTCGGCGACG ACTGCGGCCA GTCGTCGCG GTCGACGACG TCGGCGGCCA
34321 CGTACCGCAC GCGGTGCTCC TCCGGCGTGT CGCCGGGCCG GCCGTTGCGG GACACCACGA
34381 CGACCTCGGC GGCTCGTGC ACGGTGAGCA GGTGGTCCAC GAGGAGCGCG CCGAGCCCGC
34441 CCGTGCCGCC GGTGACGAGG ACGGTCCCGC CGGTGAGCGG GGAGGTTCCG GTGGCCGCGG
15 34501 CGACACGGCG CAGACGGGCC GCACGCGCTG TGCCGTGCGC GACCCGACG TGCGGCTCGT
34561 CGCCGGCGGC GAGCCCGGCC GCTATGGCGG CGGGCGTGAT CTCGTCCGCT TCGATCAGGG
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34681 CGGGATCGCC GGTACGGGTG GCCACGATGA GCCGGGATCG CGCCAGCGC GGCTCGGCGA
20 34741 GCCAGGTCTG CACGGTGGTG AGCAGGTGCG GGCCAGCTC CCGGGTCCGG GCGCCGGGCG
34801 AGGTGCCCGG GTCGCCGGGT TCCACGGCCA GGACCACGAC CGGGGGGTGC TCGCGTCCG
34861 GCACGTGCGC GAGGTACGTC CAGTCGGGGA CGGGTGACGC GGGCACGGG ACCCAGGCGA
34921 TCTCGAACAG CGCCTCGGCA TCGGGGTGCG CGGCCCGCAC GGTGAGGCTG TCGACGTCAA
34981 GGACCGGTGA GCCGTGCTCG TCCGTGGCGA CGATGCGGAC CATGTCGGGG CCGACGCGTT
25 35041 CCAGCAGCAC GCGCAGCGCG GTCGCGGCGC GCGCGTGGAT CCTCACGCC GACCAGGAGA
35101 ACGCCAGCCG GCGCCGCTCC GGGTCCGTGA AGACCGTCCC GAGGGCGTGC AGGGCCGCGT
35161 CGAGCAGCAC GGGGTGACG CCGTACCGGG CGTCGGTGAG CTGTTCCGGC AGGCGGACCG
35221 ACGCGTAGGC GCGGCCCTCC CCCGTCCACA TCGCGGTCAT GGCCCGGAAC GCGGGCCCGT
35281 ACGAGAGCGG CAGCGCGTCG TAGAAGCCGG TCAGGTGCGC CGGGTCCGGC TCGGCGGGCG
30 35341 GGCAGTCCAC GGGCTCCGCC GGACCGCCAG TGTCCACGCT CAGCGCTCCG GTGCACTGA
35401 GCGCCAGGG GCGCGTGCCG GTACGGCTGT GCAGACTCAC CGACCGCCGT CCGGACACCT
35461 CCGTTCCGAC GGTGGCTGG ATCTCCGTGT CGCCGTGCGC GTCGACCACC ACCGGCGCGA
35521 CGATGGTCAG CTCCGCGATC TCCGCGTGC CGAGCCGGGC TCCCGCTTCG GCGAGCAGTT
35581 CCACGAGCGC CGAGCCGGGC ACATGACCCG GCGCGTCCAC CTCGTGGTCG GCGAGGTCAG
35641 GCTGACGGCG TACCGAGACA CCGCGGTGGC CAGCGCGCCC TCGCCGTGCG GCGAGGTGCGA
35 35701 CCCACGAGCC GAGCAGCGGG TGGCCGGACG TTCCCGCCGG TTCCCGGTGCG ATCCAGTAGC
35761 GGTACGCGCG GAACGGGTAC GTGGGACGCG GCACACCCG ACGCGTCGCG AACGACCAGG
35821 TGACGGGCAC GCGCCGGACC CAGAGCGCGG CGAGCGACCG AGTGAAGCGG TCCAGGCGCG
35881 CCTCGCCTCG CCGCAGTGTG CCGGTGACGA CCGTATGCGC ATGCCCCGCG AGCGTGTCTT
40 35941 CCAGTGCGGT GGTGAGCACG GGATGCGCGC TGACCTCGAC GAACGCGCGG TATCCGCGGT
36001 CCGCCAGGTG GCGGTCGCG GCGGCGAACC GAACGGTGCG GCGCAGGTTG TCGTACCAGT
36061 AGGCGGCGTC CGCGGGCCGG TCCAGCCACG CCTCGTCCAC GGTGAGAGA AACGGGACGT
36121 CCGGCGTGCG CGGAGTGATG CCGGCGAGAG CGTCGAGCAG CGCGCCGCGG ATCGTTTCGA
36181 CATGCGCGGT GTGCGACGCG TAGTCGACGG CGATCCGGCG GCGCGGGGGG GTGGCGGCCA
36241 GCAGCTCCTC CACGGCGTCG GCCGCACCGG CGACAACGAT CGACGCGGGT CCGTTGACCG
45 36301 CGGCGACCTC CAGGCGCCCC GCGCACACGG CGGCGTCGAA GTCGGCGGGG GGCACCGAGA
36361 CCATGCCCGC CTGCCCCGCC AGTTCGGTGG CGACGAGTCG GCTGCGCACC GCGACGACCT
36421 TCGCGGCGTC GTCCAGGGTG AGCACCCCGG CGACGAGGC CGCGGCGACT TCGCCCTGGG
36481 AGTGGCCGAC GACCGCGGCC GGGGCGACCC CGTGCGCACG CCACAGCTCC GCCAGCGCCA
50 36541 CCATCACCGC GAACGACGCG GGTCGACAGA CATCGACCCG GTCGAACGCG GCGGCTCCGG
36601 GCGGCTGGGC GATGACGTCC AGCAGTCCC ATCCGGTGTG CCGGGCGAGC GCCGTGGCGC
36661 ACTCGCGGAG CCGCCGGGCG AACACGGGCT CCGTGGCGAG CAGTTCGGCA CCCATGCCGG
36721 CCCACTGGGA GCGCTGCCCC GGGAACGCGA ACACGACACG TGTGTCGGTG ACGTCGGCGG
36781 TTCCCGTCAC GCGCCCCGCG ACTTCGGCAC CACGGGCGAA CGCCTCCGCC TCTCGGGCCG
36841 GCACGACCGC CCGGTGGCGC ATGGCCGTCC GGGTGGTGGC GAGCGAGTGG CCGACCGCGG

- 40 -

FOCUS 40315-082701

	36901	CCGCGGCGCC	AGTGAGCGGG	GCCAGCTGTC	CCGCGACGTC	CCGCAGTCCC	TCCGGGGTCC
	36961	GGGCCGACAT	CGGCCAGACC	ACGTCTCTCG	GCACCGGCTC	GGCTTCGGGT	GCGGACACGG
	37021	GTGCGGGCGC	GGCGGGGGGC	CCGGCCTCCA	GGACGACATG	GGCGTTGGTG	CCGCTGATGC
5	37081	CGAACGACGA	GACACCCGCA	CGCCGGGCGC	GCCCGGTGAC	CGGCCACGGC	TCACTGCGGT
	37141	GCAGCAGCCG	GATGTCGCGG	TCCCAGTCTGA	CGTGCCGGGA	CGGCTCGTCG	ACGTGCAGCG
	37201	TGCGCGGCAG	GACGCCGTGC	CGCATCGCCA	TGACCATCTT	GATGACGCCG	GCGACGCCGG
	37261	CCGCGGCCCTG	GGTGTGGCCG	ATGTTTCACT	TGAGCGAGCC	GATCAGCAGC	GGATGCACGC
	37321	GTTCGCGCCC	GTAGGCCACT	TGCAGGGCCT	GGGCCTCGAC	GGGGTTCGCCG	AGACGGGTGC
	37381	CGGTGCCGTG	TGCCTCCACG	CGCTCGACGT	CACCCGGCGC	CAGGCCGGCG	TCGGCGAGCG
10	37441	CACGCTGGAT	GACGCGCTGC	TGCCGAGGCC	CGTTTCGGGGC	GGACAGCCCCG	TTTCGACGCGC
	37501	CGTCGGAGTT	GACCGCGGAG	CCGCGCACCA	GCGCCAGCAC	GGGGTGGCCG	TGGCGGGTGG
	37561	CGTCGGAGAG	CCGCTCCAGC	ACCAGGACAC	CGGCGCCCTC	GGCGAAGCTC	GTGCCGTCCG
	37621	CGGTGTCCGC	GAAGGCCTTG	GCACGGCCGT	CGGGGGCGAG	CCCGCGCTGC	CCGCGAGACT
	37681	CGACGAACCC	GGTCGTCTGC	GCCATCACCG	TGACACCGCC	GACCAGGGCG	AGCGAGCACT
15	37741	CCCCGAGCG	CAGCGACCGC	GCGGCCTGGT	GCAGCGCCAC	CAGCGACGAC	GAACACGCCG
	37801	TGTCGACGGT	GACCGACGGG	CCCTCCAGAC	CGAAGTAGTA	CGAGAGCCGC	CCGGAGAGAA
	37861	CGCTGGTTCG	CGTGCCGGTC	GCCCCGAAAC	CGCCCAGGTC	CACGCCCCGC	CCGTAGCCCT
	37921	GGGTGAACGC	GCCCATGAAT	ACGCCGGTGT	CGCTGCCCGG	GACGCTTTTCG	GGCAGGATGC
20	37981	CCGCTCGTTC	GAACGCCTCC	CACGACGCTT	CGAGGACCAG	ACGCTGCTGC	GGGTCCATCG
	38041	CCAGCGCCTC	ACGCGGGCTG	ATCCCGAAGA	ACGCGGCGTC	GAAGTCGGCG	GCGCCGGTGA
	38101	GGAAGCCGCC	GTGACGCACG	GAAACCTTGC	CGACCGCGTC	GGGGTTCGGG	TCGTAGAGCG
	38161	CGGCGAGGTC	CCAGCCGCGG	TCGGCGGGGA	ACTCGGTGAT	CGCGTCCCCG	CCGAGTCTGA
	38221	CCAGCCGCCA	CAGGTCTCTC	GGTGACCGCA	CGCCACCGGG	CATCCGGCAC	GCCATGGCCA
25	38281	CGATCGCCAG	CGGCTCGTTC	CCCGCCACCG	TCGGTGCGGG	CACTGTCTGCC	GCCGGAGCGG
	38341	CAGGGGCCGG	CTACCCCGC	CGTTCCTCAT	CCAGGCGGGC	GGCGAGCGCG	GCCGGTGTTCG
	38401	GGTGGTCGAA	GACGGCCGTC	GCGGAGAGCC	GTACCCCGCT	CGTCTCGGCG	AGGCTGTTGC
	38461	GCAACCCGAC	ACCGTGAGC	GAGTCGATGC	CGAGGTCCTT	GAACGCCGTC	GTGGGCGTGA
	38521	TCTCGGAGGC	GTCCGGGTGG	CCGAGCACGG	CGGCCGTGGC	CGCACACACG	ATGGCCAGCA
30	38581	GGTCACGATC	GCGGTCTCGG	TCGCGGTTCG	GGTTGTCTTC	CGCACGGGCG	GCGATGCGGC
	38641	GCTCGGTCCG	CTGCCGGACG	GGCTCGGTGG	GAATCGCCGC	GACCATGAAC	GGCACGTCCG
	38701	CGGCGAGGCT	CGCGTCGATG	AAGTGGGTGC	CCTCGGCCCTC	GGTGAGCGCG	CGGAACCCGT
	38761	CGCGCACCCG	CTGCCGGTTCG	GCGTCGTCAA	GTTGTCCGGT	GAGGGTCTCTG	GTGGTGTGCC
	38821	ACATGCCCCA	GGCGATGGAG	GTGGCGGGTT	GGCCGAGGGT	GTGGCGGTGG	GTGGCGAGGG
35	38881	CGTCGAGGAA	GGCGTTGGCG	GCGGCGTAGT	TTCCTTGTCC	GGGGCTGCCG	AGGACGGCGG
	38941	CGGCGCTGGA	GTAGAGGACG	AAGTGGGTGA	GGGGTTGGTT	TTGGGTGAGG	TGGTGCAGGT
	39001	GCCAGGCGGC	GTTGGCTTTG	GGGTGGAGGA	CGGTGGTGAG	GCGGTCGGGG	GTGAGGGCGT
	39061	CGAGGATGCC	GTCGTCTGAGG	GTGGCGGCGG	TGTGGAAGAC	GCGGTCGAGG	GGTTGGGGGA
	39121	TGTGGGCGAG	GGTGGTGGCG	AGTTGGTGGG	GGTCGCCGAC	GTCGCAGGGG	AGGTGGGTGC
40	39181	CGGGGGTGGT	GTCGGGGGGT	GGGGTGCGGG	AGAGGAGGTA	GGTGTGGGGG	TGGTTCAGGT
	39241	GGCGGGCGAG	GATGCCGGCG	AGGGTGCCGG	AGCCGCCGGT	GATGATGATG	GCGTGTTCGG
	39301	GGTTGAGGGG	GGTGGTGGTG	GGTGGGGTGG	TGGTGTGGAG	GGGGGTGAGG	TGGGGTCTGGT
	39361	GGAGGGTGTG	GTGGGTGAGG	CGGAGGTGGG	GGTGGTCGAG	GGTGGCGAGT	TGGGCCAGGG
	39421	GGAGGGGAGT	GTGGGGGTGG	TCGGTTTTCGA	TGAGGCGGAT	GCGGTGGGGG	TGTTCTGTTCT
	39481	GGGCGGTGCG	GGTGAGGCCG	GTGACGGTGG	CGCCGGCGGG	GTCGGTGGTG	GTGTGGACGA
45	39541	TGAGGGTGTG	GTCGGTGGTG	GTGAGGTGGT	GTTGCAGGGC	GGTCAGGACG	CGGGTGGCGC
	39601	GGGTGTGGGC	GCGGGTGGGT	ATGTCTCTCG	GGTCGTCCGG	GTGGGCGGCG	GTGATCAGGA
	39661	CGTGTCCTCT	GGGCAGGTCA	CCGTCTGTA	CCGCCTCGGC	GACCGCGAGC	CACTCCAACC
	39721	GGAGCGGGTT	CGGCCCCGAC	GGGTGTCTCG	CCCGCTCCCT	CAGCACCAGC	GAGTCCACCG
	39781	ACACGACAGG	ACGGCCATCC	GGGTCTGGCA	CGCGCACGGC	GACGCCGGCC	TCCCCCGGG
50	39841	TGAGGGCGAC	GCGCACCGCG	GCGGCCCCGG	TGGCGTTTCAG	GCGCACGGCC	GTCCTAGAGA
	39901	ACGGCAGCTC	GATCCCCCGG	CCCGCGTCTGA	GGCGCCCCGGC	GTGCAGGGCC	GCTCAGGACA
	39961	GTGCCGGATG	CACACCGAAA	CCGTCCGCCT	CGGCGGCCTG	CTCGTCTGGG	AGCGCCACCT
	40021	CGGCATACAC	GGTGTACCA	TCACGCCAGG	CAGCCCCGAA	CCCCTGGAAC	GCCGACCCGT
	40081	ACTCATAACC	GGCATCCCGC	AGTTCGTCAT	AGAACCCCGA	GACGTCGACG	GCCGCGGCGG

- 41 -

40141	TGGCCGGCGG	CCACTGCGAG	AACGGCTCAC	CGGAAGCGTT	GGAGGTATCC	GGGGTGTCCG
40201	GGGTCAGGGT	GCCGCTGGCG	TGCCGGGTCC	AGCTGCCCCG	GCCCTCGGTA	CGCGCGTGGA
40261	CGGTCACCGG	CCGCCGTCCG	GCCTCATCGG	CCCCTTCCAC	GGTCACCGAC	ACATCCACCG
5	40321	CTGCGGTAC	CGGCACCACG	AGCGGGGATT	CGATGACCAG	TTCATCCACC
	40381	CGGTCTCGTC	ACCGGCCCGG	ATGACCAGCT	CCACAAACGC	CGTACCCGGC
	40441	TGCCCCGCAC	CGCGTGATCA	GCCAGCCAGG	GATGCGTACG	CAATGAGATC
	40501	GAACAACACC	ACCACCGTCG	TGGCGGGGCA	GTGCTGTGAC	GGCGGCCAGC
	40561	CCGCCCCGGT	CAGCCCGGCC	GCGGACAGGT	CGGTGGCACC	GGCCGCCTCC
	40621	GCCTGTGCTC	GAACGCGTAG	GTGGGCAGAT	CCAGCAGCCG	CCCCGGCACC
10	40681	CCGTGCCCCA	GTCCACCCCC	GCACCCAGAG	TCCACGCCTG	CGCCAACGCC
	40741	GCTCCCAGCC	ACCGTCACCA	GTCCGCAACG	ACGCCACCGT	GCGGGCCTGT
	40801	GCAGCAGCAC	CGGATGGGCA	CTGCACTCCA	CGAACACCGA	CCCGTCCAGC
	40861	CCGCATCCAG	CGCGACAGGG	CGACGACGGT	TCCGGTACCA	GTACCCCTCA
	40921	CGGTCAACCA	GGCGCTGTCC	ACGGTCGACC	ACCACGCCAC	CGACCCGGTC
15	40981	TTCCCTTCAG	TACCTCAGCG	AGTTCGTCCT	CGATGGCCTC	CACGTGAGGC
	41041	CGTAGTCGAC	CGCGATACGA	CGCACCCGCA	CCCCATCAGC	CTCATACCGC
	41101	CCTCCACCGC	CGACGGGTCC	CCCGCCACCA	CCGTGCAAGC	CGGACCATTA
	41161	TCCACACACC	CTCGACCAGA	CCCACCTCAC	CGGCCGGCAA	CGCCACCGAA
	41221	CCCGGCCGGC	CAGCCGCGCC	GCGATCACCC	GACTGCGCAA	CGCCACCACG
20	41281	CCTCCAGGCT	GAGGGCTCCG	GCCACACACG	CCGCCGCGAT	CTCCCCCTGC
	41341	CCACAGCGTC	CGGCACGACC	CCATGCGCCT	GCCACAGCGC	GGCCAGGCTC
	41401	CCCAGCTGGC	CGGCTGGACC	ACCTCCACCC	GCTCCGCCAC	ATCCGACCGC
	41461	CCCGCACATC	CCAGCCCCTG	TGCGGCAACA	ACGCCCGCGC	ACACTCCTCC
	41521	CGAACACCGC	GGAACGGTCC	ATGAGTTCCA	CGCCCATGCC	CACCCACTGG
25	41581	CGGGGAAGAC	GAACACCGTA	CGCGGCTGAT	CCACCGCCAC	ACCCATCACC
	41641	CCAGCAGCAC	CGCACGGTGA	CCGAAGACAG	CACGCTCACG	CACCAACCCC
	41701	CGGCCACATC	CACCCACCC	CCGCGCAGAT	ACCCCTCCAG	CCGCTCCACC
	41761	GACTCACCTC	ACCACGAGCC	GACACCGGCA	ACGGCACCAA	CCCATCACCA
	41821	CACGCGACGG	CCCAGGAACA	CCCTCCAGGA	TCACGTGCGC	GTTGTAACCG
30	41881	ACGACGACAC	ACCCGCATCG	GGTGCCCGAT	CCGACTCGGG	CCACGGCCTC
	41941	GCAGCTCCAC	CGCACCGGCC	GACCACTCCA	CATGCGACGA	CGGCTCGTCC
	42001	TCTTCGGCGC	GATCCCATGC	CGCATCGCCA	TGACCATCTT	GATGACACCG
	42061	CAGCCGCTCG	CGCATGACCG	ATGTTGCACT	TGACCGAACC	GAGGTAGAGC
	42121	GGTCCTGCCC	GTAGGCCGCG	AGGACGGCCT	GCGCCTCGAT	CGGGTCGCCC
35	42181	CGGTGCCGTG	CGCCTCCACC	ACGTCCACAT	CGGCGGCGCG	CAGTCCGGCG
	42241	CCTGCCGGAT	CACGCGCTGC	TGGGCGACGC	CGTTGGGGGC	GGACAGTCCG
	42301	CGTCCTGGTT	CACCGCCGAG	CCGCGGACGA	CCGCGAGAAC	GGTGTGCCCC
	42361	CGTCGGAGAG	CCGCTCCAGC	ACGAGAACGC	CGACGCCCTC	GGCGAAGCCG
	42421	CCGCGTCGGC	GAACGCCTTG	CACCGTCCGT	CCGGGGAGAG	TCCGCGCTGC
40	42481	CCACGAGCTC	TGCGGTGTTT	GCCATGACGG	TGACACCGCC	GACCAGCGCC
	42541	CCCGGGCCCC	CAGTGCCCTG	GCCGCTGGT	GCAGGGCGAC	CAGCGACGAC
	42601	TGTCGACCGT	GACCGCCGGG	CCCTGAAGTC	CGTACACGTA	CGAGAGGCGC
	42661	CGTCGCTCTG	CGTCGCCGTG	ACACCGAGCC	CGCCCAGGTC	CCGGCCGACG
	42721	GGTTGAACGC	GCCCATGAAC	ACGCCGGTGT	CGCTCTCCCG	GAGCCTGTCC
45	42781	CGGCGTTCTC	GAACGCCCTC	CAGGAGGTCT	CCAGGATCAG	GCGCTGCTGG
	42841	CCAGCGCCTC	GTTCCGACTG	ATGCCGAAGA	ACGCGGCGTC	GAACCCGGCG
	42901	ATCCGCGCTG	GCGTGTCGTG	GAGCGGCCGG	CCGCGTCCGG	GTCCGGGTCC
	42961	CGACGTCCCA	GCCCCGGTCC	GTGGGGAAC	CGGTGATCGC	CTCGGTACCG
	43021	GCCGCCACAG	GTCCCTCCGC	GAGGCGACCC	CGCCGGGCAG	TCGGCACGCC
50	43081	TCGCGACGGG	GTCGCCGGAG	CCGAGGGTCT	GGGCGGTGCG	GGGTGCCGCT
	43141	CGGCGAGGTG	GGCGGCCAAC	GCACGCGGAG	TGGGTGGTGC	GAACCGCGTT
	43201	CCCGCAGACC	CGTCCGCGCG	GCGACGGTGT	TGGTGAAC	GACGGTGGTG
	43261	GGCCGTTCTC	GCGGAACGTG	CGGTCCGGGG	AGCAGTGTCC	GGCGCCCGGC
	43321	CGGTGGCGAC	GCTGTCGCGG	ACCAGGTCGA	GCAGTACGTC	CTCCCCGCCC

- 42 -

004046 004046 004046

	43381	CGGCGAGGCG	GTTCGCCCCAC	TCCTGTTCCG	TGGCGTCGGG	CTCGGCCGGT	CCGGTCAGTG
	43441	CGGTGAGGAT	CGGCGGCGTG	GCGCCCGCCA	TCGTGCGGCG	CCGCGCCCCG	GCGGAACCGG
	43501	TCCGGGCCAC	GATGTACGAG	CCGCCGCCCG	CGATGGCCTT	CTCGATCAGG	TCGCCGGTGA
5	43561	GCGCCGGCCG	TTCGATGCCG	GGCAGCGCGC	GGACGGTGAC	GGTGGGGAGT	CCCTCCGCGG
	43621	CCCGTGGCCG	GGTGTGGGCG	TCGGCGCCGG	CCGGGCCGTC	GAGCAGGACG	TGCACGAGCG
	43681	CGCCGGGGTT	CGCGGCTTCC	TCGGCTGCGG	TGGTCACGTG	GGTGAGGCCG	GTCTCGTCGC
	43741	GGAGCAGGCC	GGCGACGGTG	TCGGCGTCTT	CCCCGGTGAC	CAGGACCGGC	GCGTCCGGGC
	43801	CGATCGGAGG	CGGCACGGTG	AGGACCATCT	TGCCGGTGTG	CCGGGCGTGG	CTCATCCACG
10	43861	CGAACGCGTC	CCGCGCACGG	CGGATGTCCC	ACGGCTGCAC	CGGCAGCGGG	CACAGCTCAC
	43921	CGCGGTGAA	CAGGTGAGG	AGCAGTTCGA	GGATCTCCCG	CAGGCGCGCG	GGATCCACGT
	43981	CGGCCAGGTC	GAACGGCTGC	TGGGCGGCGT	GGCGGATGTC	GGTCTTGCCC	ATCTCGACGA
	44041	ACCGGCCGCC	CGGTGCGAGC	AGGCCGATGG	ACGCGTCGAG	GAGTTCACCG	GTGAGCGAGT
	44101	TGAGCACGAC	GTCGACCGGC	GGGAAGGTGT	CGCGAACGCG	GGCGCTGCGG	GAGTTCGCCA
	44161	CATGGTCGGT	GTCGAAGCCG	TCGGCGTGCA	GCAGGTGTTG	TTTGGCGGGA	CTGGCGGTGG
15	44221	CGTACACCTC	GGCGCCGAGG	TGGCGGGCGA	TCCGGGTGCG	CGCCATGCCG	ACACGGCCCC
	44281	TCGCGGCGTG	GACCAGGACC	TTCTGGCCGG	GTCGCAGCTC	GCCCGCGTCG	ACGAGGCCGT
	44341	ACCAGGCGGT	GGCGAACACG	ATGGGCACGG	ACGCGGCGAT	GGGGAACGAC	CATCCCCGTG
	44401	GGATCCGTGC	GACCAGCCGC	CGGTCCGCGA	CCACGCTGCG	CCGGAACGCG	TCCTGCACGA
20	44461	GACCGAACAC	GCGGTGCGCG	GGGGCCAGGT	CGTCGACGCC	GGGTCCGACT	TCGGTCACGA
	44521	TGCCCCGCGC	CTCCCCGCCC	ATCTCGCCCT	CGCCCGGGTA	GGTGCCGAGC	GCGATCAGCA
	44581	CGTCGCGGAA	GTTTCAGCCCC	GCGGCGCGGA	CGTCGATGCG	GACCTCGCCG	GCGGCCAGGG
	44641	GCGCGGCGGG	ACGTCGAGCG	GGGCGACGAC	GAGGTGCGCG	AGCGTTCCGG	AGGCGGGCGG
	44701	GCGCAGCGCC	CACTGGCGCG	GTCGGCAGGG	GGGTGGTGTC	CGCGCGTACC	AGCCGGGGCA
25	44761	CGTAGGCCAC	GCCGGCCCCG	AGCGCGATCT	GGGGTTCGCC	GAGCGAGGCC	GCGGCGGGGA
	44821	CGAGGTGCTC	ATCGCCGTCC	GTGTCCACCA	GCACGAACGA	TCCGGGTTCG	GCGGCCTGGC
	44881	GGCGCAGCGC	CTCGTCCCAG	AGCCGGGCCT	GGTCCGCGTC	CGGGATCTCG	GCCGGGCCGA
	44941	CGCCCACCGC	GCGGCGGGTG	ACGACCCTCC	GGCGGGGTGA	CGGGGTGCCG	GGCAGGTGCG
	45001	GCCGCTCCCA	GACCAGTTCG	CACAGCGTGG	CCTCGCCACT	GCCGGTGGCG	ACCAGATGGG
30	45061	CCGGCAGCCC	CGCGAGCCCG	GCGCGCTGGA	CCTTGCCCCA	CGCGGTGCGG	GGGATCGTGG
	45121	TGACGTGCCA	GATCTCGTCG	GGCACCTTGA	AGTAGGCGAG	CCGGCGGCGG	CACTCGGCGA
	45181	GGATCGCCTC	GGCGGGGACG	GCGGGGCCGT	CGGAAACGAC	GTAGAGCACG	GGTATGTGCG
	45241	CGAGGACGGG	GTGCGGGGCG	CCCGCCGCGT	CGGCGTCCCG	GACACCGGCC	ACCTCCTGGG
	45301	CGACGGTCTC	GATCTCCCGG	GGGTGGATGT	TCTCCCCGCC	GCGGATGATC	AGCTCCTTGA
35	45361	CCCGGCCCGT	GATCGTCACG	TGTCCGGTCT	CGGCCTGACG	TGCGAGGTCC	CCGGTCCGGT
	45421	ACCAGCCGTC	CACGAGCACC	TGGGCGGTG	CCTCCGGCTG	GGCGTGGTAG	CCGAGCATGA
	45481	GGCTCGGCCC	GCTCGCCCAC	AGCTCGCCCT	CCTCGCCGGG	TGCCACGTG	GCGCCGGACA
	45541	CCGGGTGAC	GAACCGCAGC	GACAGGCCCG	GCACGGGCAG	CCCGCACGAG	CCGGGAACCC
	45601	GCGCATCCTC	CAGGGTGTG	GCGGTGAGCG	AGCCGGTCGT	CTCGGTGACG	CCGTACGTGT
40	45661	CGAGCAGGGG	CACGCCGAAC	GTCGCCTCGA	AATCCCTGGT	GAGCGACGCC	GGCGAGGTGG
	45721	ATCCGGCGAC	CAGCGCCACG	CGCAGCGCGC	GAGCCCGCGG	CTCGCCGGAC	ACGGCGCCGA
	45781	GGAGGTAGCG	GTACATCGTC	GGCACGCCGA	CGAGCACGGT	GCTGGAGTGT	TCGGCCAGGG
	45841	CGTCGAGGAC	GTCACGCGCG	ACGAAGCCGC	CCAGGATACG	GGCGGACGCG	CCGACCGTGA
	45901	GGACGGCGAG	CAGGCAGAGG	TGGTGGCCGA	GGCTGTGGAA	CAGCGGGGCG	GGCCAGAGCA
45	45961	GTTCGTGCTC	CTCGGTCAGC	CGCCAGGACG	GCACGTGCGA	GTGCATCGCG	GACCACAGGC
	46021	CGCTGCGCTG	TGCGGAAACC	ACGCCCTTGG	GACGGCCGGT	GGTGCCGGAG	GTGTAGAGCA
	46081	TCCAGGCGGG	TTCGTCCAGG	CCGAGGTGCT	CGCGGGGCGG	GCACGGCGGC	TCGGTCCCGG
	46141	CGAGGTCCCT	GTAGGAGACG	CAGTCCGGTG	CCCGGCGCCC	GACGAGCACG	ACGGTGGCGT
	46201	CGGTGCCGGT	GCGGCGCACG	TGGTCGAGGT	GGGTTTCGTC	GGTGACCAGC	ACGGTCGCGC
50	46261	CGGAGTCCGT	CAGGAAAGTG	GCGAGTTCGG	CGTCGGCGGC	GTCCGGGTTG	AGCGGGACGG
	46321	CGACGGCGGC	GGCGCGGGCG	GCGGCGAGGT	AGACCTCGAT	GGTCTCGATC	CGGTTGCCGA
	46381	GCAGCATCGC	GACCCGGTCC	CCGCGGTGCA	GCOCGGACGC	GGCGAGGTGT	CCGGCGAGCC
	46441	GGCCGGCCCC	GAGCCGGAGT	TGCGTGTAAC	TCACGGCGCG	TTGGGAATCC	CTGTAGGCGA
	46501	TCCGGTCCGC	GCGTCGCTCG	GCATGGATGC	GGAGCAATTC	GTGCAACGCG	CGGATTGGTT
	46561	CCACACGCGC	CATGGAAACA	CCTTTCTCTC	GACCAACCGC	ACAACAGCAC	GGAACCGGCC

- 43 -

46621	ACGAGTAGAC	GCCGGCGACG	CTAGCAGCGT	TTTCCGGACC	GCCACCCCT	GAAGATCCCC
46681	CTACCGTGGC	CGGCCTCCCC	GGACGCTCAT	CTAGGGGGTT	GCACGCATAC	CGCCGTGCGT
46741	AATTGCCTTC	CTGATGACCG	ATGCCGGACG	CCAGGGAAGG	GTGGAGGCGT	TGTCCATATC
46801	TGTCACGGCG	CCGTATTGCC	GCTTCGAGAA	GACCGGATCA	CCGGACCTCG	AGGGTGACGA
5	46861	GACGGTGCTC	GGCCTGATCG	AGCACGGCAC	CGGCCACACC	GACGTGTCTG
	46921	TGCTCCCCGG	ACCGCCGTGC	ACACCACGAC	CCGTGACGAC	GAGGCGTTCA
	46981	GCACGCACAG	CGCCCTGTCT	AGTCCGGCAT	GGACAACGGC	ATCGCCTGGG
	47041	CGCGTACCTG	TTCGGTGTCT	TGCGCACCGG	CGAGAGCGGC	AGGTACGCCG
	47101	GGCCCTCTAC	ACGAACGTCT	TCCAGCTCAC	CCGGTCTGCT	GGGTATCCCC
10	47161	GACCTGGAAC	TACGTCAGCG	GTATCAACAC	GACGAACGCG	GACGGGCTGG
	47221	GGACTTCTGC	GTGGGCCGCG	CCCAGGCGCT	CGACGAGGGC	GGGATCGACC
	47281	GCCCGCGGCC	ACCGGTATCG	CGCGCCACGG	GGGCGGCATC	ACCTGCGTGT
	47341	CCGGGGCGGA	GTGCGGATCA	ACATCGAGAA	CCCCGCCGTC	CTCACGGCCC
	47401	GACGACGTAC	GGTCCGCGGC	CCCCGGTCTT	CGCACGGGCC	ACCTGGCTGG
15	47461	GGGGGGCCGG	CTGTTTCATCT	CCGCGACGGC	CGGCATCCTC	GGACACCGAA
	47521	CGGTGATGTG	ACCGGCCAGT	GCGAGGTCTG	CCTCGACAAC	ATGGCCCCGG
	47581	GGAGAACCTG	CGGCGCCACG	GCGTCCAGCG	GGGGCACGTC	CTCGCCGACG
	47641	CAAGGTCTAC	GTCCGCCGCC	GCGAGGATCT	CGATACGGTC	CGCCGGGTCT
	47701	CCTGTCGAGC	ACCGCGGCCG	TCGCCCTTTT	GCACACCGAC	ATAGCCCGCG
20	47761	CGTCGAAATC	GAAGGCATGG	TGGCGTGACA	ATACCCGGTA	AAAGGCCCGC
	47821	CTCGGCGGAT	CCGCGAAGAG	AAAGAAGAGC	GTCACCGCAC	AGCGCGGCAG
	47881	TCGTCTTTCG	CACAGCGGCG	GATCTGGTTT	CTCCAGCAAT	TGGACCCGGA
	47941	TATAATCTCC	CGCTCGTGCA	ACGCCTGCGC	GGTCTATTGG	ACGCGCCGGC
	48001	GCGCTGGCGC	TCGTCTGTCG	GCGCCACGAG	GCGTTGCGGA	CGGTGTTTCA
25	48061	GGCGAGCCCC	TCCAGCGGGT	GCTTCCCGCC	CCGGAACACC	TCCTGCGCCA
	48121	GGCAGCGAGG	AGGACGCCGC	CCGGCTCGTC	CGCGACGAGA	TCGCCGCGCC
	48181	GCCACCGGGC	CGTTGATCAG	GGCCCTGCTG	ATCCGCCTCG	GTGACGACGA
	48241	GCGGTGACCG	TGCACCATGT	CGCCGGCGAC	GGCTGGTCTG	TCGGGCTCCT
	48301	CTCGCAGCCC	ACTACACGGC	GCTGCGCGAC	ACTGCCCGCC	CTGCCGAAC
30	48361	CCGGTGACGT	ACGCCAGCTT	CGCCGCTTGG	GAGCGGCGCG	AACTCACC
	48421	GACAGGCGTC	TGGCCTACTG	GCGCGAGCAA	CTCCGGGGCG	CCCCGGCGCG
	48481	CCCACCGACC	GTCCCCGCCC	GCCGGTCCGC	GACGCGGACG	CGGGCATGGC
	48541	CCGCCGGCCG	CGCTGGCCAC	CGCGGTCTCT	ACGCTCGCGC	GCGACTCCGG
	48601	TTCATGACCC	TGCTGGCGGC	CTTCCAAGCG	GTCCTCGCCC	GGCAGGCGGG
35	48661	GTGCTGGTCT	GCACGCCCGT	GGCGAACCCT	ACGCGGGCGG	CGTACGAGGG
	48721	ATGTTCTGTC	ACACGCTCGC	GCTGCGCGGC	GACCTCTCGG	GCGATCCGTC
	48781	CTCCTCGACC	GCTGCCGGGC	CACGACCACG	GACGCGTTCT	CCCACGCCGA
	48841	GAGAACGTCA	TCGAACTCGT	CGCACCAGAA	CGCGACCTGT	CGGTCAACCC
	48901	GTGCTGTTGC	AGGTGCTGCG	GCGCGACGCG	GCGACGGCCG	CGCTGCCCGG
40	48961	GAACCGTTCC	GCACCGGACG	CTGGTTTACC	CGCTTCGACC	TCGAATTCCA
	49021	GAGCCGGGTG	GCGCGCTGAC	CGGCGAAGTG	CTCTACAGCC	GTGCGCTGTT
	49081	CGGATCACGG	GGTTGCTGGA	GGAGTTCACG	GCGGTGCTTC	AGGCGGTAC
	49141	GACGTACGGC	TGTCGCGGCT	GCCGGCCGGC	GACGCGACGG	CGGCAGCGCC
	49201	TCGAACGACA	CGGCGCGGGA	CCTGCCCGTC	GACACGCTGC	CGGGCCTGCT
45	49261	GCCGCACGCA	CCCCCGGCGC	CGTGGCCGTC	ACCGACCCGC	ACATCTCCCT
	49321	CAGCTGGACC	GGCGGGCGAA	CCGCCTCGCG	CACCTGCTCC	GCGCGCGCGG
	49381	GCGCAGCTGG	TCGGGATCTG	CGCCGATCGC	GGCGCCGACC	TGATCGTCTG
	49441	ATCCTCAAGG	CGGGCGCCGC	TTATGTGCCG	CTGGACCCCG	AACATCCTCC
	49501	GCGTTCTGTG	TGGCCGACGC	CGAGCTGACC	ACGGTGTTGG	CGCACGAGGT
50	49561	CGGTTCCCCG	ATGTGCCGCA	CGTGGTGCGC	TTGGACGACC	CGGAGCTGGA
	49621	GACGACACGG	CGCCGGACGT	CGAGCTGGAC	CGGGACAGCC	TCGCCTACGC
	49681	TCCGGGTCGA	CCGGCAGGCC	GAAGGCCCGT	CTCATGCCGG	GTGTACGCGC
	49741	CTGCTCTGGC	AGGAGCGCAC	GATGGGCCGC	GAGCCGGCCA	GCCGCACCGT
	49801	ACGCCCACGT	TCGACTACTC	GGTGCAGGAG	ATCTTTTCCG	CGCTGCTGGG

- 44 -

5 49861 GTCATCCCGC CGGACGAGGT GCGGTTTCGAC CCGCCGGGAC TCGCCCGGTG GATGGACGAA
49921 CAGGCGATTA CCCGGATCTA CGCGCCGACG GCCGTACTGC GCGCGCTGAT CGAGCACGTC
49981 GATCCGCACA GCGACCAGCT CGCCGCCCTG CCGCACCTGT GCCAGGGCGG CGAGGCGCTG
50041 ATCCTCGACG CGCGGTTGCG CGAGCTGTGC CCGCACCGGC CCCACCTGCG CGTGACAAT
50101 CACTACGGTC CGGCCGAAAG CCAGCTCATC ACCGGGTACA CGCTGCCCCG CGACCCCGAC
50161 GCGTGGCCCG CCACCGCACC GATCGGCCCG CCGATCGACA ACACCCGCAT CCATCTGCTC
50221 GACGAGGCGA TGCGGCCGGT TCCGGACGGT ATGCCGGGGC AGCTCTGCGT CGCCGGCGTC
50281 GGCCTCGCCC GTGGGTACCT GGCCCGTCCC GAGCTGACCG CCGAGCGCTG GGTGCCGGGA
50341 GATGCGGTGCG GCGAGGAGCG CATGTACCTC ACCGGCGACC TGGCCCGCCG CGCGCCCGAC
10 50401 GGCGACCTGG AATTCCTCGG CCGGATCGAC GACCAGGTCA AGATCCGCGG CATCCGCGTC
50461 GAACCGGGTG AGATCGAGAG CCTGCTCGCC GAGGACGCCC GCGTCACGCA GCGGGCGGTG
50521 TCCGTGCGCG AGGACCGGCG GGGCGAGAAG TTCCTGGCCG CGTACGTCGT ACCGGTGGCC
50581 GGCCGGCAGC GCGACGACTT CGCCGCGCTG CTGCGCGCGG GACTGGCCGC CCGGCTGCCC
50641 GCGCGGCTCG TGCCCTCCGC CGTCGTCTG GTGGAGCGAC TGCCGAGGAC CACGAGCGGC
15 50701 AAGGTGGACC GCGCGCGGCT GCGCGACCCG GAGCCGGGCC CCGGCTCGAC CGGGCGGTT
50761 ACGCCCCGCA CCGATGCCGA GCGGACGGTG TGCCGGATCT TCCAGGAGGT GCTCGACGTC
50821 CCGCGGGTGC GTGCCGACGA CGACTTCTTC ACGCTCGGCG GGCACCTCCCT GCTCGCCACC
50881 CGGGTCTGCT CCCGCATCCG CGCCGAGCTG GGTGCCGATG TCCCGCTGCG TACGCTCTTC
50941 GACGGGCGGA CGCCCGCCGC GCTCGCCGT GCGGCGGACG AGGCCGGCCC GGCGGCCCTG
20 51001 CCCCCGATCG CGCCCTCCGC GGAGAACGGG CCGGCCCCCC TCACCGCGGC ACAGGAACAG
51061 ATGCTGCACT CGCACGGCTC GCTGCTCGCC GCGCCCTCCT ACACGGTCGC CCCGTACGGG
51121 TTCCGGCTGC GCGGGCCACT CGACCGCGAA GCGCTCGACG CCGCACTGAC CCGGATCGCC
51181 GCGCGCCACG AGCCGCTGCG GACCGGGTTC CCGGATCGGG AACAGGTCGT CCGGCCGCCC
51241 GTCGCGGTGC GCGCCGAGGT GGTTCCGGTG CCGGTGCGCG ACGTCGACGC CGCGGTCCGG
25 51301 GTCGCCCACC GGGAGCTGAC CCGGCCGTTT GACCTCGTGA ACGGGTCGTT GCTGCGTGCC
51361 GTGCTGCTGC CGCTGGGCGC CGAGGATCAC GTGCTGCTGC TGATGCTGCA CCACCTCGCC
51421 GGTGACGGAT GGTCTTTCGA CCTCTGGTC CGGGAGTTGT CCGGGACGCA ACCCGACCTT
51481 CCGGTGTCCT ACACGGACGT GGCCCGGTGG GAACGGAGTC CCGCCGTGAT CGCGGCCAGG
51541 GAGAACGACC GGGCTACTG GCGCCGGCGG CTGGGGGGCG CCACCGCGCC GGAGCTGCCC
30 51601 GCGGTCGCGC CCGGCGGGG ACCGACGGG CCGGCGTTCC TGTGGACGCT CAAGGACACC
51661 CCGGTCTTGG CGGCACGCC GGTGCGGAG GCCACGACG CGACGTTGCA CGAAACCGTG
51721 CTCGGCGCCT TCGCCCTGGT CGTGGCGGAG ACCGCCGACA CCGACGACGT GCTCGTCGCG
51781 ACGCCGTTTC CGGACCGGGG GTACGCGGG ACCGACCACC TCATCGGCTT CTTGCGGAAG
51841 GTCCTCGCGC TGCGCCTCGA CCTCGGCGGC ACGCCGTCGT TCCCGAGGT TCCGCGCGG
35 51901 GTGCACACCG CGATGGTGG CGCGCACGCC CACCAGGCGG TGCCCTACTC CGCGCTGCGC
51961 GCCGAGGACC CCGCGCTGCC GCGGGCCCC GTGTGCTTCC AGCTCATCAG CGCGCTCAGC
52021 GCGGAAGTGC GGCTGCCCG CATGCACACC GAGCCGTTCC CCGTCGTCGC CGAGACCGTC
52081 GACGAGATGA CCGGCGAACT GTCGATCAAC CTCTTCGACG ACGGTGCGAC CGTCTCCGGC
52141 GCGGTGGTCC ACGATGCCGC GCTGCTCGAC CGTGCCACCG TCGACGATT GCTACCCCGG
40 52201 GTGGAGGCGA CGCTGCGTGC CGCCGCGGG GACCTACCG TACGCGTAC CGGTTACGTG
52261 GAAAGCGAGT AGCCATGCCC GAGCAGGACA AGACAGTCGA GTACCTTCGC TGGGCGACCG
52321 CGGAAGTCCA GAAGACCCGT GCGGAAGTC CCGCGCACAG CGAGCCGTTG GCGATCGTGG
52381 GGATGGCCTG CCGGCTGCCC GGCGGGGTG CGTCGCCGGA GGACCTGTGG CAGTTGCTGG
52441 AGTCCGGTGG CGACGGCATC ACCGCGTTCC CCACGGACCG GGGCTGGGAG ACCACCGCCG
45 52501 ACGGTCGCGG CGGCTTCCTC ACCGGGGCGG CCGGCTTCGA CGCGGCGTTC TTCGGCATCA
52561 GCGGCGCGCA GGCGCTGGCG ATGGACCCGC AGCAGCGCCT GGCCCTGGAG ACCTCGTGGG
52621 AGGCGTTCGA GCACGCGGGC ATCGATCCGC AGACGCTGCG GGGCAGTGAC ACGGGGGTGT
52681 TCCTCGGCGC GTTCTTCCAG GGGTACGGCA TCGGCGCCGA CTTGACGGT TACGGCACCA
52741 CAGGACATTA CACGAGCGTG CTCTCCGGCC GCCTCGCGTA CTTCTACGGT CTGGAGGGTC
50 52801 CCGGCGTCAC GGTGACACG CGGTGTTGCT CGTCGCTGGT GGCGCTGCAC CAGGCCGGGC
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52921 CGCCGGCGGG GTTCGCGGAC TTCTCCGAGC AGGGCGGCCT GGCCCCGAC GCGCGCTGCA
52981 AGGCCTTCGC GGAAGCGGCT GACGGCACCG GTTTCGCCGA GGGGTCCGGC GTCCTGATCG
53041 TCGAGAAGCT CTCCGACGCC GAGCGCAACG GCCACCGCGT GCTGGCGGTC GTCCGGGGTT

- 45 -

53101	CCGCCGTCAA	CCAGGACGGT	GCCTCCAACG	GGCTGTCCGC	GCCGAACGGG	CCGTCGCAGG
53161	AGCGGGTGAT	CCGGCAGGCC	CTGGCCAACG	CCGGACTCAC	CCCGGCGGAC	GTGGACGCCG
53221	TCGAGGCCCA	CGGCACCGGC	ACCAGGCTGG	GCGACCCCAT	CGAGGCACAG	GCCGTGCTGG
53281	CCACCTACGG	GCAGGGGCGC	GACACCCCTG	TGCTGCTGGG	CTCGCTGAAG	TCCAACATCG
53341	GCCACACCCA	GGCCGCCGCG	GGCGTCGCCC	GTGTCATCAA	GATGGTCCCT	GCCATGCGGC
53401	ACGGCACCCT	GCCCCGCACC	CTGCACGTGG	ACACGCCGTC	CTCGCACGTC	GACTGGACGG
53461	CCGGCGCCGT	CGAACTCCTC	ACCGACGCCC	GGCCCTGGCC	CGAAACCGAC	CGCCACGGC
53521	GCGCCGGTGT	CTCCTCCTTC	GGCGTCAGCG	GCACCAACGC	CCACATCATC	CTCGAAAGCC
53581	ACCCCGACCC	GGCCCCCGAA	CCCGCCCCGG	CACCCGACAC	CGGACCGCTG	CCGCTGCTGC
53641	TCTCGGCCCG	CACCCCGCAG	GCACTCGACG	CACAGGTACA	CCGCCTGCGC	GCGTTCCTCG
53701	ACGACAACCC	CGGCGCGGAC	CGGGTCGCCC	TCGCGCAGAC	ACTCGCCCCG	CGCACCCAGT
53761	TCGAGCACCG	CGCCGTGCTG	CTCGCGGACA	CGCTCATCAC	CGTGAGCCCG	AACGCCGGCC
53821	GCGGACCGGT	GGTCTTCGTC	TACTCGGGGG	AAAGCACGCT	GCACCCGCAC	ACCGGGCGGC
53881	AACTCGCGTC	CACCTACCCC	GTGTTCCGCC	AAGCGTGCGG	CGAGGCCCTC	GACCACCTCG
53941	ACCCACCCCA	GGGCCCGGCC	ACGCACTTCG	CCCACCAGAC	CGCGCTCACC	GCGCTCCTGC
54001	GGTCTTGGGG	CATCACCCCG	CACGCGGTCA	TCGGCCACTC	CCTCGGTGAG	ATCACCCCGG
54061	CGCACGCCGC	CGGTGTCTTG	TCCCTGAGGG	ACGCGGGCGC	GCTCCTCACC	ACCCGCACCC
54121	GCCTGATGGA	CCAAGTCCCG	TCGGGCGGCG	CGATGGTCAC	CGTCTTGACC	AGCGAGGAAA
54181	AGGCACGCCA	GGTGCTGCGG	CCGGGCGTGG	AGATCGCCGC	CGTCAACGGC	CCCCACTCCC
54241	TCGTGCTGTC	CGGGGACGAG	GAAGCCGTAC	TCGAAGCCGC	CCGGCAGCTC	GGCATCCACC
54301	ACCGCCTGCC	GACCCGCCAC	GCCGGCCACT	CCGAGCGCAT	GCAGCCACTC	GTGCCCCCCC
54361	TCCTCGACGT	CGCCCGGACC	CTGACGTACC	ACCAGCCCCA	CACCGCCATC	CCCGGCGACC
54421	CCACCACCGC	CGAATACTGG	GCGCACCAGG	TCCGCGACCA	AGTACGTTTC	CAGGCGCACA
54481	CCGAGCAGTA	CCCGGGGCGG	ACGTTCCTCG	AGATCGGCCC	CAACCAGGAC	CTCTCGCCGC
54541	TCGTGACCGG	CGTTGCCGCC	CAGACCCGTA	CGCCCCGACG	GGTGCGGGCG	CTGCACACCG
54601	CGCTCGCGCA	GCTCCACGTC	CGCGGCGTCG	CGATCGACTG	GACGCTCGTC	CTCGGCGGGG
54661	ACCGCGCGCC	CGTCACGCTG	CCACAGTATC	CGTTCCAGCA	CAAGGACTAC	TGGCTGCGGC
54721	CCACCTCCCG	GGCCGATGTG	ACCGGCGCGG	GGCAGGAGCA	GGTGGCGCAC	CCGCTGCTCG
54781	GCGCCGCGGT	CGCGCTGCCC	GGCACGGGCG	GAGTCGTCTT	GACCGGCCGC	CTGTGCTTGG
54841	CCTCCCATCC	GTGGCTCGGC	GAGCACGCGG	TCGACGGCAC	CGTGCTCCTG	CCCGGCGCGG
54901	CCTTCCTCGA	ACTCGCGGCG	CGCGCCGGCG	ACGAGGTCGG	CTGCGACCTG	CTGCACGAAC
54961	TCGTATCATGA	GACGCCGCTC	GTGCTGCCCC	CGACCGGCGG	TGTGGCGGTC	TCCGTGAGAA
55021	TCGCCGAACC	CGACGACACG	GGGCGGCGGG	CGGTACCCGT	CCACGCGCGG	GCCGACGGCT
55081	CGGGCCTGTG	GACCCGACAC	GCCGGCGGAT	TCCTCGGCAC	GGCACCGGCA	CCGGCCACGG
55141	CCACGGACCC	GGCACCCCTG	CCGCCCGCGG	AAGCCGGACC	GGTCGACGTC	GCCGACGTCT
55201	ACGACCGGTT	CGAGGACATC	GGGTACTCCT	ACGGACCGGG	CTTCCGGGGG	CTGCGGGCCG
55261	CCTGGCGCGC	CGGCGACACC	GTGTACGCCG	AGGTGCGGCT	CCCCGACGAG	CAGAGCGCCG
55321	ACGCCGCCCG	TTTCACGCTG	CACCCCGCGC	TGCTCGACGC	CGCGTTCCAG	GCCGGCGCGC
55381	TGGCCGCGCT	CGACGCACCC	GGCGGGGCGG	CCCGACTGCC	GTTCTCGTTC	CAGGACGTCC
55441	GCATCCACGC	GGCCGGGGCG	ACGCGGCTGC	GGGTACGCGT	CGGCCGCGAC	GGCGAGCGCA
55501	GCACCGTCCG	CATGACCGGC	CCGGACGGGC	AGCTGGTGCC	CGTGGTCCGT	GCCGTGCTGT
55561	CGCGCCCGTA	CGCGGAAGGC	TCCGGTGACG	GCCTGCTGCG	CCCGGTCTGG	ACCGAGCTGC
55621	CGATGCCCGT	CCCGTCCCGG	GACGATCCGC	GCGTGAGAGT	CCTCGGCGCC	GACCCGGGCG
55681	ACGGCGACGT	TCCGGCGGCC	ACCGGGGAGC	TGACCGCCCC	CGTCCTCGGC	GCGCTCCAGC
55741	GCCACCTGTC	CGCCGCCGAG	GACACCACCT	TGGTGGTACG	GACCGGCACC	GGCCCGGCCG
55801	CTGCCGCCGC	CGCGGGTCTG	GTCCGCTCGG	CGCAGGCGGA	GAACCCCGGC	CGCGTCGTGC
55861	TCGTGAGGCG	GTCCCCGGAC	ACCTCGGTGG	AGCTGCTCGC	CGCGTGCGCC	GCGCTGGACG
55921	AACCCGACGT	GGCCGTCCGG	GACGGCGTGC	TCTTCGCGCC	GCGGCTGGTC	CGGATGTCCG
55981	ACCCCGCGCA	CGGCCCGCTG	TCCCTGATCG	ACGGCGACTG	GCTGCTCACC	CGGTCCGCCT
56041	CCGGCACGTT	GCACGACGTC	CGCGTCCATG	CCGACGACAC	GCCCCGCGCG	GCGCTCGAAG
56101	CCGGCGAGGT	CCGCATCGAC	GTCCGCGCGG	CCGGACTGAA	CTTCCGCGAT	GTGCTGATCG
56161	CGCTCGGGAC	GTACACCGGG	GCCACGGCCA	TGGGCGGGCA	GGCCGCGGGC	GTGCTGATGG
56221	AGACCGGGCC	CGGCGTGGAC	GACCTGTCCC	CCGGCGACCG	GGTGTTCGGC	CTGACCCGGG
56281	GCGGCATCGG	CCCGACGGCC	GTCACCGACC	GGCGCTGGCT	GGCCCGGATC	CCCGACGGCT

56341	GGAGCTTCAC	CACGGCGGCG	TCCGTCCCGA	TCGTGTTCGC	GACCGCGTGG	TACGGCCTGG
56401	TCGACCTCGG	CACACTGCGC	GCCGGCGAGA	AGGTCCTCGT	CCACGCGGCC	ACCGGCGGTG
56461	TCGGCATGGC	CGCCGCACAG	ATCGCCCGCC	ACCTGGGCGC	CGAGCTCTAC	GCCACCGCCA
56521	GTACCGGCAA	GCAGCACGTC	CTGCGCGCCG	CCGGGCTGCC	CGACACGCAC	ATCGCCGACT
56581	CTCGGACGAC	CGCGTTCCGG	ACCGCTTTCC	CGCGCATGGA	CGTCGTCTTG	AACGCGCTGA
56641	CCGGCGAGTT	CATCGACGCG	TCGCTCGACC	TGCTGGACGC	CGACGGCCGG	TTCTGTCGAGA
56701	TGGGCCGCGC	CGAGCTGCGC	GACCCGGCCG	CGATCGTCCC	CGCCTACCTG	CCGTTCGACC
56761	TGCTGGACGC	GGGCGCCGAC	CGCATCGGCG	AGATCCTGGG	CGAACTGCTC	CGGCTGTTTCG
56821	ACGCGGGCGC	GCTGGAGCCG	CTGCCGGTCC	GTGCCTGGGA	CGTCCGGCAG	GCACGCGACG
56881	CGCTCGGCTG	GATGAGCCCG	GCCCGCCACA	TCGGCAAGAA	CGTCTGACG	CTGCCCCGGC
56941	CGCTCGACCC	GGAGGGCGCC	GTCGTCTCTA	CCGGCGGCTC	CGGCACGCTC	GCCGGCATCC
57001	TCGCCCCGCA	CCTGCGCGAA	CGGCATGTCT	ACCTGCTGTC	CCGGACGGCA	CCGCCCCGAGG
57061	GGACGCCCCG	CGTCCACCTG	CCCTGCGACG	TCGGTGACCG	GGACCAGCTG	GCGGCGGCCC
57121	TGGAGCGGGT	GGACCGGCCG	ATCACCGCCG	TGGTGACCTT	CGCCGGTGCG	CTGGACGACG
57181	GCACCGTCGC	GTGCTCACC	CCCGAGCGTT	TCGACACGGT	GCTGCGCCCC	AAGGCCGACG
57241	GCGCCTGGTA	CCTGCACGAG	CTGACGAAGG	AGCAGGACCT	CGCCGCGTTC	GTGCTCTACT
57301	CGTCGGCCGC	CGGCGTGCTC	GGCAACGCCG	GCCAGGGCAA	CTACGTCGCC	GCGAACGCGT
57361	TCCTCGACGC	GCTCGCCGAG	CTGCGCCACG	GTTCCGGGCT	GCCGGCCCTC	TCCATCGCCT
57421	GGGGGCTCTG	GGAGGACGTG	AGCGGGCTCA	CCGCGGCGCT	CGGCGAAGCC	GACCGGGACC
57481	GGATGCGGCG	CAGCGGTTTC	CGGGCCATCA	CCGCGCAACA	GGGCATGCAC	CTGTACGAGG
57541	CGGCCGGCCG	CACCGGAAGT	CCCGTGGTGG	TCGCGGCGGC	GCTCGACGAC	GCGCCGGACG
57601	TGCCGCTGCT	GCGCGGCCTG	CGGCGGACGA	CCGTCCGGCG	GGCCGCCGTC	CGGGAGTGTT
57661	CGTCCGCCGA	CCGGCTCGCC	GCGCTGACCG	GCGACGAGCT	CGCCGAAGCG	CTGCTGACGC
57721	TCGTCCGGGA	GAGCACCGCC	GCCGTGCTCG	GCCACGTGGG	TGGCGAGGAC	ATCCCCGCGA
57781	CGGCGGCGTT	CAAGGACCTC	GGCATCGACT	CGCTCACCGC	GGTCCAGCTG	CGCAACGCCC
57841	TCACCGAGGC	GACCGGTGTG	CGGCTGAACG	CCACGGCGGT	CTTCGACTTC	CCGACCCCGC
57901	ACGTGCTCGC	CGGGAAGCTC	GGCGACGAAC	TGACCGGCAC	CCGCGCGCCC	GTCGTGCCCC
57961	GGACCGCGGC	CACGGCCGGT	GCGCACGACG	AGCCGCTGGC	GATCGTGGGA	ATGGCCTGCC
58021	GGCTGCCCCG	CGGGTGCCCG	TCACCCGAGG	AGCTGTGGCA	CCTCGTGGCA	TCCGGCACCG
58081	ACGCCATCAC	GGAGTTCCCC	ACGGACCGCG	GCTGGGACGT	CGACGCGATC	TACGACCCGG
58141	ACCCCGACGC	GATCGGCAAG	ACCTTCGTCC	GGCACGGTGG	CTTCCTCACC	GGCGCGACAG
58201	GCTTCGACGC	GGCGTTCTTC	GGCATCAGCC	CGCGCGAGGC	CCTCGCGATC	GACCCGAGC
58261	AGCGGGTGCT	CCTGGAGACG	TCGTGGGAGG	CGTTTCAAAG	CGCCGGCATC	ACCCCGGACT
58321	CGACCCGCGG	CAGCGACACC	GGCGTGTTTC	TCGGCGCCTT	CTCCTACGGT	TACGGCACCG
58381	GTGCGGACAC	CGACGGCTTC	GGCGCGACCG	GCTCGCAGAC	CAGTGTGCTC	TCCGGCCGGC
58441	TGTCGTACTT	CTACGGTCTG	GAGGGTCCGG	CGGTACGGT	CGACACGGCG	TGTTCTGTCG
58501	CGCTGGTGGC	GCTGCACCAG	GCCGGGCAGT	CGCTGCGCTC	CGGCGAATGC	TCGCTCGCCC
58561	TGGTCGGCGG	CGTCACGGTG	ATGGCGTCTC	CCGGCGGCTT	CGTGGAGTTC	TCCCGGCAGC
58621	GCGGCCTCGC	GCCGGACGGC	CGGGCGAAGG	CGTTCCGGCG	GGGTGCGGAC	GGCACGAGCT
58681	TCGCCGAGGG	TGCCGGTGTG	CTGATCGTCG	AGAGGCTCTC	CGACGCCGAA	CGCAACGGTC
58741	ACACCGTCCT	GGCGGTCGTC	CGTGGTTCGG	CGGTCAACCA	GGATGGTGCC	TCCAACGGGC
58801	TGTGCGCGCC	GAACGGGCCG	TCGCAGGAGC	GGGTGATCCG	GCAGGCCCTG	GCCAACGCCG
58861	GGCTCACCCC	GGCGGACGTG	GACGCCGTCT	AGGCCACCG	CACCGGCACC	AGGCTGGGCG
58921	ACCCCATCGA	GGCACAGGCG	GTAATGGCCA	CCTACGGACA	GGAGCGCGCC	ACCCCCCTGC
58981	TGCTGGGCTC	GCTGAAGTCC	AACATCGGCC	ACGCCCAGGC	CGCGTCCGGC	GTCGCCGGCA
59041	TCATCAAGAT	GGTGACGGCC	CTCCGGCACG	GGGAGCTGCC	GCCGACGCTG	CACGCCGACG
59101	AGCCGTCGCC	GCACGTCGAC	TGGACGGCCG	GCGCCGTCTG	ACTGCTGACG	TCGGCCCGGC
59161	CGTGCGCCGA	GACCGACCGG	CCACGGCGTG	CCGCCGTCTC	CTCGTTCCGG	GTGAGCGGCA
59221	CCAACGCCCA	CGTCATCCTG	GAGGCGCGAC	CGGTAACGGA	GACGCCCGCG	GCATCGCCTT
59281	CCGGTGACCT	TCCCCTGCTG	GTGTCGGCAC	GCTACCGGGA	AGCGCTCGAC	GTCAGATATC
59341	GCCGACTGCG	CGCCTACCTG	GACACCACCC	CGGACGTCGA	CCGGGTGGCC	GTGGCACAGA
59401	CGCTGGCCCC	GCGCACACAC	TTCGCCCACC	GCGCCGTGCT	GCTCGGTGAC	ACCGTCATCA
59461	CCACACCCCC	CGCGGACCGG	CCCGACGAAC	TCGTCTTCGT	CTACTCCGGC	CAGGGCACCC
59521	AGCATCCCGC	GATGGGCGAG	CAGCTCGCCG	CCGCCCATCC	CGTGTTCGCC	GACGCCTGGC

- 47 -

59581 ATGAAGCGCT CCGCCGCTT GACAACCCCG ACCCCACGA CCCCACGCAC AGCCAGCATG
59641 TGCTCTTCGC CCACCAGGCG GCGTTCACCG CCCTCCTGCG GTCCTGGGGC ATCACCCTCGC
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5 59821 CACCCGGTGC CATGGTCACC GTACTGACCA GCGAAGAGAA GGCACGCCAG GCGTTGCGGC
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59941 ACGCCGTGCT CACCGTCGCC GGGCAGCTCG GCATCCACCA CCGCCTGCCC GCCCGCACG
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60061 TCCGCTACCA CCCTCCCCAC ACCTCCATTC CGAACGACCC CACCACCGCT GAGTACTGGG
10 60121 CCGAGCAGGT CCGCAAGCCC GTGCTGTTCC ACGCCACGC GCAGCAGTAC CCGGACGCCG
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15 60421 CCGACGCGGG CCACCCCGTG CTGGGCTCCG GTATCGCCCT CGCCGGGTG CCGGGCCGGG
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45 62161 GCGAGGACTT CGTCATGGCC GCCGCGATGG ACCCGGCACA GCCGATGACC GGCTCCGTAC
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62761 CCCCCGCAA GACCTACGTC CGGCACGGCG GCTTCTCTCG CGAGGCGGCC GGCTTCGATG

- 48 -

	62821	CCGCGTTCTT	CGGCATCAGC	CCGCGCGAGG	CACGGGCCAT	GGACCCGCAG	CAGCGCGTCA
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	62941	GCAGCGACAC	CGGCGTGTTT	ATGGGCGCGT	TCTCCCATGG	GTACGGCGCC	GGCGTCGACC
5	63001	TGGGCGGGTT	CGGCGCCACC	GCCACGCAGA	ACAGCGTGCT	CTCCGGCCCG	TTGTCTGACT
	63061	TCTTCGGCAT	GGAGGGCCCC	GCCGTACCCG	TCGACACCGC	CTGCTCGTCG	TCGCTGGTCG
	63121	CCCTGCACCA	GGCGGCACAG	GCGCTGCGGA	CTGGAGAATG	CTCGCTGGCG	CTCGCCGGCG
	63181	GTGTCACGGT	GATGCCCACC	CCGCTGGGCT	ACGTCGAGTT	CTGCCGCCAG	CGGGGACTCG
	63241	CCCCCGACGG	CCGTTGCCAG	GCCTTCGCGG	AAGGCGCCGA	CGGCACGAGC	TTCTCGGAGG
10	63301	GCGCCGGCGT	TCTTGTGCTG	GAGCGGCTCT	CCGACGCCGA	GCGCAACGGA	CACACCGTCC
	63361	TCCGCTCCTC	CGCTCCTCC	GCCGTCAACC	AGGACGGCGC	CTCCAACGGC	ATCTCCGCAC
	63421	CCAACGGCCC	CTCCAGCAG	CGCGTCATCC	GCCAGGCCCT	CGACAAGGCC	GGGCTCGCCC
	63481	CCGCCGACGT	GGACGTGGTG	GAGGCCACG	CACCCGGAAC	CCCGCTGGG	GACCCGATCG
	63541	AGGCACAGGC	CATCATCGCG	ACCTACGGCC	AGGACCGCGA	CACACCGCTC	TACCTCGGTT
15	63601	CGGTCAAGTC	GAACATCGGA	CACACCCAGA	CCACCGCCGG	TGTCGCCGGC	GTCATCAAGA
	63661	TGGTCATGGC	GATGCGCCAC	GGCATCGCGC	CGAAGACACT	GCACGTGGAC	GAGCCGTCGT
	63721	CGCATGTGGA	CTGGACCGAG	GGTGCGGTGG	AACTGCTCAC	CGAGGCGAGG	CCGTGGCCCC
	63781	ACGCGGGACG	CCCGCGCCGC	GCGGGCGTGT	CGTCGCTCGG	TATCAGCGGT	ACGAACGCCC
20	63841	ACGTGATCCT	TGAGGGTGTT	CCCGGGCCGT	CGCGTGTGGA	GCCGTCTGTT	GACGGGTTGG
	63901	TGCCGTTGCC	GGTGTCGGCT	CGGAGTGAGG	CGAGTCTGCG	GGGGCAGGTG	GAGCGGCTGG
	63961	AGGGGTATCT	GCGCGGGAGT	GTGGATGTGG	CCGCGGTCGC	GCAGGGGTTG	GTGCGTGAGC
	64021	GTGCTGTCTT	CGGTACCCGT	GCGGTACTGC	TGGGTGATGC	CCGGGTGATG	GGTGTGGCGG
	64081	TGGATCAGCC	GCGTACGGTG	TTCGTCTTTC	CCGGGCAGGG	TGCTCAGTGG	GTGGGCATGG
	64141	GTGTGGAGTT	GATGGACCGT	TCTGCGGTGT	TCGCGGCTCG	TATGGAGGAG	TGTGCGCGGG
25	64201	CGTTGTTGCC	GCACACGGGC	TGGGATGTGC	GGGAGATGTT	GGCGCGGCCG	GATGTGGCGG
	64261	AGCGGGTGGA	GGTGGTCCAG	CCGGCCAGCT	GGGCGGTCGC	GGTCAGCCTG	GCCGCACTGT
	64321	GGCAGGCCCA	CGGGGTGCTA	CCCGACGCGG	TGATCGGACA	CTCCCAGGGC	GAGATCGCGG
	64381	CGGCGTGCGT	GGCCGGGGCC	CTCAGCCTTG	AGGACGCCGC	CCGCGTGGTG	GCCTTGCGCA
	64441	GCCAGGTCAT	CGCGGCGCGA	CTGGCCGGGC	GGGGAGCGAT	GGCTTCGGTG	GCATTGCCCG
30	64501	CCGGTGAGGT	CGGTCTGGTC	GAGGGCGTGT	GGATCGCGGC	GCGTAACGGC	CCCGCCTCGA
	64561	CAGTCGTGGC	CGGCGAGCCG	TCGGCGGTGG	AGGACGTGGT	GACGCGGTAT	GAGACCGAAG
	64621	GCGTGCGAGT	GCGTCGTATC	GCCGTCGACT	ACGCCTCCCA	CACGCCCCAC	GTGGAAGCCA
	64681	TCGAGGACGA	ACTCGCTGAG	GTAAGTGAAG	GAGTTGCAGG	GAAGGCCCGC	TACGTTGGCT
	64741	GGTGGTCGAC	CGTGGACAGC	GCCTGGGTGA	CCGAGCCGGT	GGATGAGAGT	TACGTTGATC
35	64801	GGAACCTGCG	TCGCCCCGTC	GCGCTGGACG	CGGCGGTGGC	GGAGCTGGAC	GGGTCCGTGT
	64861	TCGTGGAGTG	CAGCGCCCAT	CCGGTGCTGC	TGCCGGCGAT	GGAACAGGCC	CACACGGTGG
	64921	CGTCGTTGCG	CACCGGTGAC	GGCGGCTGGG	AGCGATGGCT	GACGGCGTTG	GCGCAGGCGT
	64981	GGACCCTGGG	CGCGGCAGTG	GACTGGGACA	CGGTGGTTCG	ACCGGTGCCA	GGGCGGCTGC
	65041	TCGATCTGCC	CACCTACGCG	TTCGAGCGCC	GGCGCTACTG	GCTGGAAGCG	GCCGGTGCCA
40	65101	CCGACCTGTC	CGCGGCCGGG	CTGACAGGGG	CAGCACATCC	CATGCTGGCC	GCCATCACGG
	65161	CACTACCCGC	CGACGACGGT	GGTGTGTTTC	TCACCGGCCG	GATCTCGTTG	GCGACGCATC
	65221	CCTGGCTGGC	TGATCACGCG	GTGCGGGGCA	CGGTCTGCTG	GCCGGGCACG	GCCTTTGTGG
	65281	AGCTGGTCAT	CCGGGCCCGT	GACGAGACCG	GTTGCGGGAT	AGTGGATGAA	CTGGTCATCG
	65341	AATCCCCCCT	CGTGGTGCCG	GCGACCGCAG	CCGTGGATCT	GTCGGTGACC	GTGGAAGGAG
45	65401	CTGACGAGGC	CGGACGGCGG	CGAGTGACCG	TCCACGCCCG	CACCGAAGGC	ACCGGCAGCT
	65461	GGACCCGGCA	CGCCAGCGGC	ACCCTGACCC	CCGACACCCC	CGACACCCCC	AACGCTTCCG
	65521	GTGTTGTGCG	TGCGGAGCCG	TTCTCGCAGT	GGCCACCTGC	CACTGCCCGC	GCCGTCGACA
	65581	CTCTGGAGTT	CTACTTGCGC	CTGGACGCGC	TGGGCTACCG	GTTGCGACCC	ATGTTCCGCG
	65641	GAATGCGGGC	TGCCTGGCGT	GATGTGACCA	CCGTGTACGC	CGAGGTGCGC	CTCCCCGAGG
50	65701	ACCGTGCCGC	CGACGCGGAC	GATTTTCGGCA	TGCACCCGGC	GCTGCTCGAC	GCGGCCTTGC
	65761	AGAGCGGCAG	CCTGCTCATG	CTGGAATCGG	ACGCGGAGCA	GAGCGTGCAA	CTGCCGTTCT
	65821	CCTGGCACGG	CGTCCGGTTC	CACGCGACCG	GCGCGACCAT	GCTGCGGGTG	GCGGCTGTAC
	65881	CGGGCCCGGA	CGGCCTCCGG	CTGCATGCCG	CGGACAGCGG	GAACCGTCCC	GTCGCGACGA
	65941	TCGACGCGCT	CGTGACCCGG	TCCCCGGAAG	CGGACCTCGC	GCCCCCGGAT	CCGATGCTGC
	66001	GGGTGCGGTG	GGCCCCGGTG	CCCGTACCTG	CCGGGGCCGG	TCCGTCCGAC	GCGGACGTGC

- 49 -

5 66061 TGACGCTGCG CGGCGACGAC GCCGACCCGC TCGGGGAGAC CCGGGACCTG ACCACCCGTG
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- 50 -

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69421	ATGCCACGGC	CGCGGCCCGT	GTCGCGGGCG	TCATCAAGAT	GGTGCAGGCG	ATCGGCGCGG
69481	GCACGATGCC	GCGGACGCTG	CATGTGGAGG	AGCCCTCGCC	CGCCGTCGAC	TGGAGCACCG
5 69541	GACAGGTGTC	CCTGCTCGGC	TCCAACCGGC	CCTGGCCGGA	CGACGAGCGT	CCGCGCCGGG
69601	CGGCCGTCTC	CGCGTTCGGG	CTCAGCGGGA	CGAACGCGCA	CGTCATCCTG	GAACAGCACC
69661	GTCCGGCGCC	CGTGCGCTCC	CAGCCGCCCC	GGCCGCCCCG	TGAGGAGTCC	CAGCCGCTGC
69721	CGTGGGTGCT	CTCCGCGCGG	ACTCCGGCCG	CGCTGCGGGC	CCAGGCGGGC	CGGCTGCGCG
69781	ACCACCTCGC	GGCGGCACCG	GACGCGGATC	CGTTGGACAT	CGGGTACGCG	CTGGCCACCA
10 69841	GCCGCGCCCA	GTTCGCCCAC	CGTGCCGCGG	TCGTCGCCAC	CACCCCGGAC	GGATTCCGTG
69901	CCGCGCTCGA	CGGCCTCGCG	GACGGCGCGG	AGGCGCCCGG	AGTCGTCAAC	GGGACCGCTC
69961	AGGAGCGGCG	CGTCGCCTTC	CTCTTCGACG	GCCAGGGCGC	CCAGCGCGCC	GGAATGGGGC
70021	GCGAGCTCCA	CCGCCGGTTC	CCCGCTCTTC	CCGCCGCGTG	GGACGAGGTC	TCCGACGCGT
70081	TCGGCAAGCA	CCTCAAGCAC	TCCCCACAGG	ACGTCTACCA	CGGCGAACAC	GGCGCTCTCG
15 70141	CCCATGACAC	CCTGTACGCC	CAGGCCGGCC	TGTTACAGCT	CGAAGTGGCG	CTGCTGCGGC
70201	TGCTGGAGCA	CTGGGGGGTG	CGGCCGGACG	TGCTCGTCGG	GCACTCCGTC	GGCGAGGTGA
70261	CCGCGGCGTA	CGCGGCGGGG	GTGCTCACCC	TGGCGGACGC	GACGGAGTTG	ATCGTGGCCC
70321	GGGGGCGGGC	GCTGCGGGCG	CTGCCGCCCG	GGGCGATGCT	CGCCGTCGAC	GGAAGCCCGG
70381	CGGAGGTCGG	CGCCCGCACG	GATCTGGACA	TCGCCGCGGT	CAACGGCCCG	TCCGCCGTGG
20 70441	TGCTCGCCGG	TTCGCCGGAC	GATGTGGCGG	CGTTCGAACG	GGAGTGGTCG	GCGGCCGGGC
70501	GGCGCACGAA	ACGGCTCGAC	GTCGGGCACG	CGTTCCACTC	CCGGCACGTC	GACGGTGC GC
70561	TCGACGGCTT	CCGTACGGTG	CTGGAGTCGC	TCGCGTTCGG	CGCGGCGCGG	CTGCCGTTGG
70621	TGTCCACGAC	GACGGGCCGG	GACGCCGCGG	ACGACCTCAT	AACGCCCGCG	CACTGGCTGC
70681	GCCATGCGCG	TCGGCCGGTG	CTGTTCTCGG	ATGCCGTCCG	GGAGCTGGCC	GACCGCGGCG
25 70741	TCACCACGTT	CGTGGCCGTC	GGCCCCCTCC	GCTCCCTGGC	GTCCGGCCGCG	GCGGAGAGCG
70801	CCGGGGAGGA	CGCCCGGACC	TACCACGCGG	TGCTGCGCGC	CCGGACCGGT	GAGGAGACCG
70861	CGGCGCTGAC	CGCCCTCGCC	GAGCTGCACG	CCCACGGCGT	CCCGGTTCGAC	CTGGCCGCGG
70921	TACTGGCCGG	TGGCCGGCCA	GTGGACCTTC	CCGTGTACGC	GTTCCAGCAC	CGTTCCTACT
70981	GGCTGGCCCC	GGCCGTGGCG	GGGGCGCCCG	CCACCGTGGC	GGACACCGGG	GGTCCGGCGG
30 71041	AGTCCGAGCC	GGAGGACCTC	ACCGTCCGCC	AGATCGTCCG	TCGGCGCACC	GCGGCGCTGC
71101	TCGGCGTCAC	GGACCCCGCC	GACGTCGATG	CGGAAGCGAC	GTTCTTCGCG	CTCGGTTTCG
71161	ACTCACTGGC	GGTGACGCGG	CTGCGCAACC	AGCTCGCCTC	GGCAACCGGG	CTGGACCTGC
71221	CGGCGGCCGT	CCTGTTCGAC	CACGACACCC	CGGCCGCGCT	CACCGCGTTC	CTCCAGGACC
71281	GGATCGAGGC	CGGCCAGGAC	CGGATCGAGG	CCGGCGAGGA	CGACGACGCG	CCCACCGTGC
35 71341	TCTCGCTCCT	GGAGGAGATG	GAGTCGCTCG	ACGCCGCGGA	CATCGCGGCG	ACGCCGGCCC
71401	CGGAGCGTGC	GGCCATCGCC	GATCTGCTCG	ACAAGCTCGC	CCATACCTGG	AAGGACTACC
71461	GATGAGCACC	GATACGCACG	AGGGAACGCC	GCCCGCCGCG	CGCTGCCCAT	TCGCGATCCA
71521	GGACGGTCAC	CGCGCCATCC	TGGAGAGCGG	CACGGTGGGT	TCGTTTCGACC	TGTTCCGGCT
71581	CAAGCACTGG	CTGGTCGCCG	CCGCCGAGGA	CGTCAAGCTG	GTCACCAACG	ATCCGCGGTT
40 71641	CAGCTCGGCC	GCGCCGTCCG	AGATGCTGCC	CGACCGGCGG	CCCGGTGGT	TCTCCGGGAT
71701	GGACTCACCG	GAGCACAACC	GCTACCGGCA	GAAGATCGCG	GGGGACTTCA	CACTGCGCGC
71761	GGCGCGCAAG	CGGGAGGACT	TCGTGCGCCG	GGCCGCCGAC	GCCTGCCTGG	ACGACATCGA
71821	GGCCGCGGGA	CCCGGCACCG	ACCTCATCCC	CGGGTACGCC	AAGCGGCTGC	CCTCCCTCGT
71881	CATCAACGCG	CTGTACGGGC	TCACCCCTGA	GGAGGGGGCC	GTGCTGGAGG	CACGGATGCG
45 71941	CGACATCACC	GGCTCGGCCG	ATCTGGACAG	CGTCAAGACG	CTGACCGACG	ACTTCTTCGG
72001	GCACGCGCTG	CGGCTGGTCC	GCGCGAAGCG	TGACGAGCGG	GGCGAGGACC	TGCTGCACCG
72061	GCTGGCCTCG	GCCGACGACG	GCGAGATCTC	GCTCAGCGAC	GACGAGGCGA	CGGGCGTGTT
72121	CGCAGCGCTG	CTGTTTCGCC	GCCACGACTC	GGTGACGACG	ATGGTCGGCT	ACTGCCTCTA
72181	CGCACTGCTC	AGCCACCCCG	AGCAGCAGGC	GGCGCTGCGC	GCGCGCCCGG	AGCTGGTCTA
50 72241	CAACGCGGTC	GAGGAGATGC	TCCGTTTCCT	GCCCGTCAAC	CAGATGGGCG	TACCGCGCGT
72301	CTGTGTTCGAG	GACGTCGATG	TGCGGGGCGT	GCGCATCCGT	GCGGGCGACA	ACGTGATCCC
72361	GCTCTACTCG	ACGGCCAACC	GCGGACCCGA	GGTGTTCCTG	CAGCCCGACA	CCTTCGATGT
72421	GACGCGCCCG	CTGGAGGGCA	ACTTCGCGTT	GCGCCACGGC	ATTCACAAGT	GTCCCGGCCA
72481	GCACATCGCC	CGGGTGCTCA	TCAAGGTCGC	CTGCCTGCGG	TTGTTTCGAGC	GTTTCCCGGA

- 51 -

72541	CGTCCGGCTG	GCCGGCGACG	TGCCGATGAA	CGAGGGGCTC	GGGCTGTTCA	GCCCGGCCGA
72601	GCTGCGGGTC	ACCTGGGGGG	CGGCATGAGT	CACCCGGTGG	AGACGTTGCG	GTTGCCGAAC
72661	GGGACGACGG	TCGCGCACAT	CAACGCGGGC	GAGGCGCAGT	TCCTCTACCG	GGAGATCTTC
72721	ACCCAGCGCT	GCTACCTGCG	CCACGGTGTC	GACCTGCGCC	CGGGGGACGT	GGTGTTCGAC
5	72781	GTCGGCGCGA	ACATCGGCAT	GTTACAGCTT	TTCGCGCATC	TGGAGTGTCC
	72841	GTGCACGCCT	TCGAGCCCGC	GCCCGTGCCG	TTCGCGGCGC	TGCGGGCGAA
	72901	CACGGCATCC	CGGGCCAGGC	GGACCAAGTC	GCGGTCTCCG	ACAGCTCCGG
	72961	ATGACCTTCT	ATCCCGACGC	CACGCTGATG	TCCGGTTTCC	ACGCGGATGC
	73021	ACGGAGCTGT	TGCGCACGCT	CGGCCTCAAC	GGCGGCTACA	CCGCCGAGGA
10	73081	ATGCTCGCGC	AACTGCCCCG	CGTCAGCGAG	GAGATCGAAA	CCCCTGTGGT
	73141	GACGTCATCG	CGGAGCGCGG	TATCGAGGCC	ATCGGCCTGC	TGAAGGTCGA
	73201	AGCGAACGGC	AGGTCTTCGC	CGGCCTCGAG	GACACCGACT	GGCCCCGTAT
	73261	GTCGCGGAGG	TCCACGACAT	CGACGGCGCG	CTCGAGGAGG	TCGTACAGCT
	73321	CATGGCTTCA	CCGTGGTTCG	CGAGCAGGAA	CCGCTGTTTC	CCGGCACGGG
15	73381	GTCGCGCGCG	GGCGGGTGGC	CGGCTGAGCG	CCGTGCGGGC	CGCGGCCGTC
	73441	GCCGCGGTGC	GGACGGCGGC	TCAGCCGGCG	TCGGACAGTT	CCTTGGGCAG
	73501	CCCTTCACCC	CCAGCTTGCG	GAACACGTTG	GTGAGGTGCT	GTTCCACCGT
	73561	ACGAACAGCT	GGCTGGCGAT	CTCCTTGTTG	GTGCGCCCGA	CCGCGGCGTG
20	73621	CGCCGCTCCG	CCTCGGTCAG	CGATGTGATC	CGCTGCGCCG	GCGTCACGTC
	73681	TCCGCGTCCG	AGGACTCCCC	ACCGAGCCGC	CGGAGGAGCG	GCACGGCTCC
	73741	GCGAGGTGCC	GTGCGCGGCG	GAACAGTCCC	CGCGCACGGC	TGTGCCGCCG
	73801	CACGCTTCGC	CCATGTGCGC	GAGGACGCGG	GCCAGCTCGT	ACTGGTCGCG
	73861	AGCAGATCGG	CGGCCTCGTC	GAGCAGTTCG	ATCCGCTTGG	CCGGCGGACT
	73921	TGCACCCGCA	GCGTCATCAC	CCGCGCCCGG	GACCCCATCG	GCCGGGACAG
25	73981	ATGAGCCTCA	GCCCCCTCGT	ACGGCCGCGG	CCGAGCAGCA	GAAGCGCTTC
	74041	ACCCGCCACA	GGGCCAGGCC	CGGCACGTCG	ACGGACCAGC	GTGCGATCCG
	74101	TCCCGGAACG	CGTTGTACGC	CGCCCGGTAC	CGCCCGGCCG	CGAGATGGTG
	74161	GCCCAGACCA	TGTGACATCC	GAAGAGGCTG	TCGGAGGTCT	CCTCCGGCAA
30	74221	AGCCACCGCT	CCGCCCCGGT	AGTCGGATCG	CGGCGGCCAC	GGTGCTGCTC
	74281	AGCGGCAATG	CGGCGGCCAT	CCCCCAGGAG	GGCACGACCC	GGGGGGCGAG
	74341	CCGCATTGCA	CGGCGGCGGT	CAGGTCGCCG	CGGCGCAGCG	CGGCCTCGGC
	74401	GCGTGGAACG	CCTCGTCGCG	CGGGGTCCGC	ATGTTGTTCG	CACCGGCCAG
	74461	CAGGACTGGA	CGGCATCGGT	GTCCTCGGCG	TAGAGCAGGG	CCAGCAACGC
35	74521	GTGGTCCGGT	CCGTGCTGAC	CCGGGAGTGC	TGGAGCACGT	ACTCGGCTTT
	74581	TGTTCCGGAC	AGCCGCGCAG	CGCGTTGCTC	AGGGCCTTGT	CGGCGACGGC
	74641	ACGGCTCCGG	AAAACGAGGC	GACCTCGTCC	TCGGCCGGCG	GATCGGCCGG
	74701	TCGGCCGCGC	CGGGATAGAT	CAGCGCGAGG	GACAGGTCCG	CGACGCGCAG
	74761	CCCTGCTCGC	TCGGGGCGGC	GGAGCGCTGG	GCCGCCAGGA	CCTCGGCGGC
40	74821	CGCCCGTCCA	TCGCCAGCCA	GCAGGCGAGC	GACACGGCGT	GCTCGCTGGA
	74881	TCCCGCGACG	CGGTGAGCAG	CTCGGGCACA	TGCCGGCCCG	ATCTGGCGGG
	74941	CGCTCGATGG	CGGCGGTGTC	GACGCGCAGT	GCGGCGTGGA	CGGCGGGGTC
	75001	CGGTAGGCGA	ACTCCAGGTA	GGTGACGGCC	TCGTGAGAGT	CGCCGCGCAG
	75061	CGCGCGGCGT	CGGTGAACAG	CCCGGCGACC	TCGGCGCCGT	GCACCCGGCC
45	75121	TGGTGGCGGG	CGAGCACCTT	GCTGGCCACG	CCGCGGTCCC	GCAGCAGTTC
	75181	TCGTGCAGGC	CACGCCGCTC	GGCGGCGGAG	AGGTCGTGCA	GTACGACGGA
	75241	GGGTGCGGGA	ACCGCCCTTC	CCGAGCAGC	CGCCCCTCGA	CCAGCTGTTC
	75301	TCGACCGCCT	CGGTGTGAGG	GCCGGTCATC	CGCTGGACGA	GGGTGAGTTC
	75361	CCGAGCACGG	CGGAAGCTCG	GGCGACGCTC	AGCGCGGCCG	GGCCGCAACG
	75421	CCGAGGTAGG	CGAGCCGGTA	CGCCCCCCCC	CGGACCACTT	CCAGGCACCC
50	75481	GTCCGTGCCT	CCCGGATGTC	GTCGATCAGG	CCGTGGCCCG	GGAGCAGGTT
	75541	GCCCGGAACG	CCTGGGCCAC	CACGTCGTCG	TGCGCGTCCT	GGCCGAGGTG
	75601	AGTTCGGTGG	TCTGCGCCTC	GGTGAGCGGG	CGCAGCGCGA	TCTCCTGGTA
	75661	CTCAGCAGTG	CCGCCCCGAA	TTGGGAGTGG	GCGGGCGTCG	GCCGGAGCAG
	75721	ACGATGGCGA	CACGGGCCCC	GCTGATGCGG	CGCGCGAGGT	GGAGCAGGCA

- 52 -

	75781	GGCGCGTCGG	CGTGGTGCAC	GTCGTGATG	CCGATCAGTA	CGGGCCGCTC	CGCGGCGAGC
	75841	GTCAGCACCG	TGCGGGTGAG	TTCGGTCCCC	AGGCGGTTGT	CGACGTCGGC	CGGCAGGTTT
	75901	TCGCACGATG	CCGTCAGCCG	GACCAGCTCC	GGTGTCCGGG	CGGCCAGCTC	GGGCTGGTCG
	75961	AGGAGCTGGC	CGAGCATGCC	GTACGGCAGG	GCCCCGCTCCT	CCATGGAGCA	CACCGCGCGA
5	76021	AGGGTGACGA	AGCCGGCCTT	GGCCGCGGCG	GCGTCGAGGA	GTTCCGGTCTT	GCCGCGAGGCG
	76081	ATCGGCCCGG	TGACGGCGGC	GACGACGCCC	CGCCCCCCCC	CCGCTCGGGT	GAGCGCCCCG
	76141	TGGAGGGAAC	CGAACTCGTC	ATCGCGGGCG	ATCAGGTCTG	GGGGAGATAA	GCGCGCTATC
	76201	ACGAATGGAA	CTACCTCGCG	ACCGTCGTGG	AAACCCATAG	GCATCACATG	GCTTGTGTGAT
	76261	CTGTACGGCT	GTGATTCAGC	CTGGCGGGAT	GCTGTGCTAC	AGATGGGAAG	ATGTGATCTA
10	76321	GGGCCGTGCC	GTTCCCTCAG	GAGCCGACCG	CCCCCGGCGC	CACCCGCGCT	ACCCCTGGG
	76381	CCACCAGCTC	GGCGACCCGC	TCCTGGTGGT	CGACGAGGTA	GAAGTGCCCG	CCGGGAAGA
	76441	CCTCCACCGT	GGTCGGCGCG	GTCGTGTGCC	CGGCCAGGC	GTGGGCCTGC	TCCACCGTCG
	76501	TCTTCGGATC	GTCGTACCG	ATGCACACCG	TGATCGGCGT	CTCCAGCGGC	GGCGCGGGCT
	76561	CCCACCGGTA	CGTCTCCGCC	GCGTAGTAGT	CCGCCGCAA	CGGCGCCAGG	ATCAGCGCGC
15	76621	GCATTTCTGTC	GTCCGCCATC	ACATCGGCGC	TCGTCCCGCC	GAGGCCGATG	ACCGCCGCCA
	76681	GCAGCTCGTC	GTCGGACGCG	AGGTGGTCCT	GGTCGGCGCG	CGGCTGCGAC	GGCGCCCGCC
	76741	GGCCCGAGAC	GATCAGGTGC	GCCACCGGGA	GCCGCTGGGC	CAGCTCGAAC	GCGAGTGTGCG
	76801	CGCCCATGCT	GTGGCCGAAC	AGCACCAGCG	GACGGTCCAG	CCCCGGCTTC	AACGCCTCGG
	76861	CCACGAGGCC	GGCGAGAACA	CGCAGGTGCG	GCACCGCCTC	CTCGTCGCGG	CGGTCTTGCG
20	76921	GGCCGGGGTA	CTGCACGGCG	TACACGTCCG	CCACCGGGGC	GAGCGCACGG	GCCAGCGGAA
	76981	GGTAGAACGT	CGCCGATCCG	CCGGCGTGGG	GCAGCAGCAC	CACCCGTACC	GGGGCCTCGG
	77041	GCGTGGGGAA	GAAGTGCCGC	AGCCAGAGTT	CCGAGCTCAC	CGCACCCCTC	CGGCCGCGAC
	77101	CTGGGGAGCC	CGGAACCGGG	TGATCTCGGC	CAAGTGCTTC	TCCCGCATCT	CCGGGTGCGT
	77161	CACGCCCCAT	CCCTCCTCCG	GCGCCAGACA	GAGGACGCCG	ACTTTGCCGT	TGTGCACATT
25	77221	GCGATGCACA	TCGCGCACCG	CCGACCCGAC	GTCGTGAGC	GGGTAGGTCA	CCGACAGCGT
	77281	CGGGTGCACC	ATCCCCTTGC	AGATCAGGCG	GTTGCGCTCC	CACGCCTCAC	GATAGTTCGC
	77341	GAAGTGGGTA	CCGATGATCC	GCTTCACGGA	CATCCACAGG	TACCGATTGT	CAAAGGCGTG
	77401	CTCGTATCCC	GAGGTTGACG	CGCAGGTGAC	GATCGTGCCA	CCCCGACGTG	TCACGTAGAC
	77461	ACTCGCGCCG	AACGTCGCGC	GCCCCGGGTG	CTCGAACACG	ATGTCGGGAT	CGTCACCGCC
30	77521	GGTCAGCTCC	CGGATC				

Those of skill in the art will recognize that, due to the degenerate nature of the genetic code, a variety of DNA compounds differing in their nucleotide sequences can be used to encode a given amino acid sequence of the invention. The native DNA sequence encoding the FK-520 PKS of *Streptomyces hygroscopicus* is shown herein merely to illustrate a preferred embodiment of the invention, and the present invention includes DNA compounds of any sequence that encode the amino acid sequences of the polypeptides and proteins of the invention. In similar fashion, a polypeptide can typically tolerate one or more amino acid substitutions, deletions, and insertions in its amino acid sequence without loss or significant loss of a desired activity. The present invention includes such polypeptides with alternate amino acid sequences, and the amino acid sequences shown merely illustrate preferred embodiments of the invention.

- 53 -

5 The recombinant nucleic acids, proteins, and peptides of the invention are many and diverse. To facilitate an understanding of the invention and the diverse compounds and methods provided thereby, the following general description of the FK-520 PKS genes and modules of the PKS proteins encoded thereby is provided. This general description is followed by a more detailed description of the various domains and modules of the FK-520 PKS contained in and encoded by the compounds of the invention. In this description, reference to a heterologous PKS refers to any PKS other than the FK-520 PKS. Unless otherwise indicated, reference to a PKS includes reference to a portion of a PKS. Moreover, reference to a domain, module, or PKS includes
10 reference to the nucleic acids encoding the same and vice-versa, because the methods and reagents of the invention provide or enable one to prepare proteins and the nucleic acids that encode them.

15 The FK-520 PKS is composed of three proteins encoded by three genes designated *fkfA*, *fkfB*, and *fkfC*. The *fkfA* ORF encodes extender modules 7 - 10 of the PKS. The *fkfB* ORF encodes the loading module (the CoA ligase) and extender modules 1 - 4 of the PKS. The *fkfC* ORF encodes extender modules 5 - 6 of the PKS. The *fkfP* ORF encodes the NRPS that attaches the pipecolic acid and cyclizes the FK-520 polyketide.

20 The loading module of the FK-520 PKS includes a CoA ligase, an ER domain, and an ACP domain. The starter building block or unit for FK-520 is believed to be a dihydroxycyclohexene carboxylic acid, which is derived from shikimate. The recombinant DNA compounds of the invention that encode the loading module of the FK-520 PKS and the corresponding polypeptides encoded thereby are useful for a variety of methods and in a variety of compounds. In one embodiment, a DNA compound
25 comprising a sequence that encodes the FK-520 loading module is inserted into a DNA compound that comprises the coding sequence for a heterologous PKS. The resulting construct, in which the coding sequence for the loading module of the heterologous PKS is replaced by the coding sequence for the FK-520 loading module, provides a novel PKS coding sequence. Examples of heterologous PKS coding sequences include the
30 rapamycin, FK-506, rifamycin, and avermectin PKS coding sequences. In another

- 54 -

embodiment, a DNA compound comprising a sequence that encodes the FK-520 loading module is inserted into a DNA compound that comprises the coding sequence for the FK-520 PKS or a recombinant FK-520 PKS that produces an FK-520 derivative.

In another embodiment, a portion of the loading module coding sequence is
5 utilized in conjunction with a heterologous coding sequence. In this embodiment, the invention provides, for example, either replacing the CoA ligase with a different CoA ligase, deleting the ER, or replacing the ER with a different ER. In addition, or alternatively, the ACP can be replaced by another ACP. In similar fashion, the corresponding domains in another loading or extender module can be replaced by one or
10 more domains of the FK-520 PKS. The resulting heterologous loading module coding sequence can be utilized in conjunction with a coding sequence for a PKS that synthesizes FK-520, an FK-520 derivative, or another polyketide.

The first extender module of the FK-520 PKS includes a KS domain, an AT
domain specific for methylmalonyl CoA, a DH domain, a KR domain, and an ACP
15 domain. The recombinant DNA compounds of the invention that encode the first extender module of the FK-520 PKS and the corresponding polypeptides encoded thereby are useful for a variety of applications. In one embodiment, a DNA compound comprising a sequence that encodes the FK-520 first extender module is inserted into a DNA compound that comprises the coding sequence for a heterologous PKS. The
20 resulting construct, in which the coding sequence for a module of the heterologous PKS is either replaced by that for the first extender module of the FK-520 PKS or the latter is merely added to coding sequences for modules of the heterologous PKS, provides a novel PKS coding sequence. In another embodiment, a DNA compound comprising a sequence that encodes the first extender module of the FK-520 PKS is inserted into a DNA
25 compound that comprises the remainder of the coding sequence for the FK-520 PKS or a recombinant FK-520 PKS that produces an FK-520 derivative.

In another embodiment, all or only a portion of the first extender module coding sequence is utilized in conjunction with other PKS coding sequences to create a hybrid module. In this embodiment, the invention provides, for example, either replacing the
30 methylmalonyl CoA specific AT with a malonyl CoA, ethylmalonyl CoA, or 2-

- 55 -

hydroxymalonyl CoA specific AT; deleting either the DH or KR or both; replacing the DH or KR or both with another DH or KR; and/or inserting an ER. In replacing or inserting KR, DH, and ER domains, it is often beneficial to replace the existing KR, DH, and ER domains with the complete set of domains desired from another module. Thus, if one desires to insert an ER domain, one may simply replace the existing KR and DH domains with a KR, DH, and ER set of domains from a module containing such domains. In addition, the KS and/or ACP can be replaced with another KS and/or ACP. In each of these replacements or insertions, the heterologous KS, AT, DH, KR, ER, or ACP coding sequence can originate from a coding sequence for another module of the FK-520 PKS, from a gene for a PKS that produces a polyketide other than FK-520, or from chemical synthesis. The resulting heterologous first extender module coding sequence can be utilized in conjunction with a coding sequence for a PKS that synthesizes FK-520, an FK-520 derivative, or another polyketide. In similar fashion, the corresponding domains in a module of a heterologous PKS can be replaced by one or more domains of the first extender module of the FK-520 PKS.

In an illustrative embodiment of this aspect of the invention, the invention provides recombinant PKSs and recombinant DNA compounds and vectors that encode such PKSs in which the KS domain of the first extender module has been inactivated. Such constructs are especially useful when placed in translational reading frame with the remaining modules and domains of an FK-520 or FK-520 derivative PKS. The utility of these constructs is that host cells expressing, or cell free extracts containing, the PKS encoded thereby can be fed or supplied with N-acylcysteamine thioesters of novel precursor molecules to prepare FK-520 derivatives. See U.S. patent application Serial No. 60/117,384, filed 27 Jan. 1999, and PCT patent publication Nos. US97/02358 and US99/03986, each of which is incorporated herein by reference.

The second extender module of the FK-520 PKS includes a KS, an AT specific for methylmalonyl CoA, a KR, an inactive DH, and an ACP. The recombinant DNA compounds of the invention that encode the second extender module of the FK-520 PKS and the corresponding polypeptides encoded thereby are useful for a variety of applications. In one embodiment, a DNA compound comprising a sequence that encodes

- 56 -

the FK-520 second extender module is inserted into a DNA compound that comprises the coding sequence for a heterologous PKS. The resulting construct, in which the coding sequence for a module of the heterologous PKS is either replaced by that for the second extender module of the FK-520 PKS or the latter is merely added to coding sequences for the modules of the heterologous PKS, provides a novel PKS coding sequence. In another embodiment, a DNA compound comprising a sequence that encodes the second extender module of the FK-520 PKS is inserted into a DNA compound that comprises the coding sequence for the remainder of the FK-520 PKS or a recombinant FK-520 PKS that produces an FK-520 derivative.

In another embodiment, all or a portion of the second extender module coding sequence is utilized in conjunction with other PKS coding sequences to create a hybrid module. In this embodiment, the invention provides, for example, either replacing the methylmalonyl CoA specific AT with a malonyl CoA, ethylmalonyl CoA, or 2-hydroxymalonyl CoA specific AT; deleting the KR and/or the inactive DH; replacing the KR with another KR; and/or inserting an active DH or an active DH and an ER. In addition, the KS and/or ACP can be replaced with another KS and/or ACP. In each of these replacements or insertions, the heterologous KS, AT, DH, KR, ER, or ACP coding sequence can originate from a coding sequence for another module of the FK-520 PKS, from a coding sequence for a PKS that produces a polyketide other than FK-520, or from chemical synthesis. The resulting heterologous second extender module coding sequence can be utilized in conjunction with a coding sequence from a PKS that synthesizes FK-520, an FK-520 derivative, or another polyketide. In similar fashion, the corresponding domains in a module of a heterologous PKS can be replaced by one or more domains of the second extender module of the FK-520 PKS.

The third extender module of the FK-520 PKS includes a KS, an AT specific for malonyl CoA, a KR, an inactive DH, and an ACP. The recombinant DNA compounds of the invention that encode the third extender module of the FK-520 PKS and the corresponding polypeptides encoded thereby are useful for a variety of applications. In one embodiment, a DNA compound comprising a sequence that encodes the FK-520 third extender module is inserted into a DNA compound that comprises the coding

- 57 -

sequence for a heterologous PKS. The resulting construct, in which the coding sequence for a module of the heterologous PKS is either replaced by that for the third extender module of the FK-520 PKS or the latter is merely added to coding sequences for the modules of the heterologous PKS, provides a novel PKS coding sequence. In another embodiment, a DNA compound comprising a sequence that encodes the third extender module of the FK-520 PKS is inserted into a DNA compound that comprises the coding sequence for the remainder of the FK-520 PKS or a recombinant FK-520 PKS that produces an FK-520 derivative.

In another embodiment, all or a portion of the third extender module coding sequence is utilized in conjunction with other PKS coding sequences to create a hybrid module. In this embodiment, the invention provides, for example, either replacing the malonyl CoA specific AT with a methylmalonyl CoA, ethylmalonyl CoA, or 2-hydroxymalonyl CoA specific AT; deleting the KR and/or the inactive DH; replacing the KR with another KR; and/or inserting an active DH or an active DH and an ER. In addition, the KS and/or ACP can be replaced with another KS and/or ACP. In each of these replacements or insertions, the heterologous KS, AT, DH, KR, ER, or ACP coding sequence can originate from a coding sequence for another module of the FK-520 PKS, from a coding sequence for a PKS that produces a polyketide other than FK-520, or from chemical synthesis. The resulting heterologous third extender module coding sequence can be utilized in conjunction with a coding sequence from a PKS that synthesizes FK-520, an FK-520 derivative, or another polyketide. In similar fashion, the corresponding domains in a module of a heterologous PKS can be replaced by one or more domains of the third extender module of the FK-520 PKS.

The fourth extender module of the FK-520 PKS includes a KS, an AT that binds ethylmalonyl CoA, an inactive DH, and an ACP. The recombinant DNA compounds of the invention that encode the fourth extender module of the FK-520 PKS and the corresponding polypeptides encoded thereby are useful for a variety of applications. In one embodiment, a DNA compound comprising a sequence that encodes the FK-520 fourth extender module is inserted into a DNA compound that comprises the coding sequence for a heterologous PKS. The resulting construct, in which the coding sequence

- 58 -

for a module of the heterologous PKS is either replaced by that for the fourth extender module of the FK-520 PKS or the latter is merely added to coding sequences for the modules of the heterologous PKS, provides a novel PKS coding sequence. In another embodiment, a DNA compound comprising a sequence that encodes the fourth extender
5 module of the FK-520 PKS is inserted into a DNA compound that comprises the remainder of the coding sequence for the FK-520 PKS or a recombinant FK-520 PKS that produces an FK-520 derivative.

In another embodiment, a portion of the fourth extender module coding sequence is utilized in conjunction with other PKS coding sequences to create a hybrid module. In
10 this embodiment, the invention provides, for example, either replacing the ethylmalonyl CoA specific AT with a malonyl CoA, methylmalonyl CoA, or 2-hydroxymalonyl CoA specific AT; and/or deleting the inactive DH, inserting a KR, a KR and an active DH, or a KR, an active DH, and an ER. In addition, the KS and/or ACP can be replaced with another KS and/or ACP. In each of these replacements or insertions, the heterologous KS,
15 AT, DH, KR, ER, or ACP coding sequence can originate from a coding sequence for another module of the FK-520 PKS, a PKS for a polyketide other than FK-520, or from chemical synthesis. The resulting heterologous fourth extender module coding sequence can be utilized in conjunction with a coding sequence for a PKS that synthesizes FK-520, an FK-520 derivative, or another polyketide. In similar fashion, the corresponding
20 domains in a module of a heterologous PKS can be replaced by one or more domains of the fourth extender module of the FK-520 PKS.

As illustrative examples, the present invention provides recombinant genes, vectors, and host cells that result from the conversion of the FK-506 PKS to an FK-520 PKS and vice-versa. In one embodiment, the invention provides a recombinant set of FK-
25 506 PKS genes but in which the coding sequences for the fourth extender module or at least those for the AT domain in the fourth extender module have been replaced by those for the AT domain of the fourth extender module of the FK-520 PKS. This recombinant PKS can be used to produce FK-520 in recombinant host cells. In another embodiment, the invention provides a recombinant set of FK-520 PKS genes but in which the coding
30 sequences for the fourth extender module or at least those for the AT domain in the fourth

- 59 -

extender module have been replaced by those for the AT domain of the fourth extender module of the FK-506 PKS. This recombinant PKS can be used to produce FK-506 in recombinant host cells.

Other examples of hybrid PKS enzymes of the invention include those in which
5 the AT domain of module 4 has been replaced with a malonyl specific AT domain to provide a PKS that produces 21-desethyl-FK520 or with a methylmalonyl specific AT domain to provide a PKS that produces 21-desethyl-21-methyl-FK520. Another hybrid PKS of the invention is prepared by replacing the AT and inactive KR domain of FK-520
10 extender module 4 with a methylmalonyl specific AT and an active KR domain, such as, for example, from module 2 of the DEBS or oleandolide PKS enzymes, to produce 21-desethyl-21-methyl-22-desoxo-22-hydroxy-FK520. The compounds produced by these hybrid PKS enzymes are neurotrophins.

The fifth extender module of the FK-520 PKS includes a KS, an AT that binds methylmalonyl CoA, a DH, a KR, and an ACP. The recombinant DNA compounds of the
15 invention that encode the fifth extender module of the FK-520 PKS and the corresponding polypeptides encoded thereby are useful for a variety of applications. In one embodiment, a DNA compound comprising a sequence that encodes the FK-520 fifth extender module is inserted into a DNA compound that comprises the coding sequence for a heterologous PKS. The resulting construct, in which the coding sequence for a
20 module of the heterologous PKS is either replaced by that for the fifth extender module of the FK-520 PKS or the latter is merely added to coding sequences for the modules of the heterologous PKS, provides a novel PKS. In another embodiment, a DNA compound comprising a sequence that encodes the fifth extender module of the FK-520 PKS is inserted into a DNA compound that comprises the coding sequence for the FK-520 PKS
25 or a recombinant FK-520 PKS that produces an FK-520 derivative.

In another embodiment, a portion of the fifth extender module coding sequence is utilized in conjunction with other PKS coding sequences to create a hybrid module. In this embodiment, the invention provides, for example, either replacing the methylmalonyl CoA specific AT with a malonyl CoA, ethylmalonyl CoA, or 2-hydroxymalonyl CoA
30 specific AT; deleting any one or both of the DH and KR; replacing any one or both of the

- 60 -

DH and KR with either a KR and/or DH; and/or inserting an ER. In addition, the KS and/or ACP can be replaced with another KS and/or ACP. In each of these replacements or insertions, the heterologous KS, AT, DH, KR, ER, or ACP coding sequence can originate from a coding sequence for another module of the FK-520 PKS, from a coding
5 sequence for a PKS that produces a polyketide other than FK-520, or from chemical synthesis. The resulting heterologous fifth extender module coding sequence can be utilized in conjunction with a coding sequence for a PKS that synthesizes FK-520, an FK-520 derivative, or another polyketide. In similar fashion, the corresponding domains in a module of a heterologous PKS can be replaced by one or more domains of the fifth
10 extender module of the FK-520 PKS.

In an illustrative embodiment, the present invention provides a set of recombinant FK-520 PKS genes in which the coding sequences for the DH domain of the fifth extender module have been deleted or mutated to render the DH non-functional. In one such mutated gene, the KR and DH coding sequences are replaced with those encoding
15 only a KR domain from another PKS gene. The resulting PKS genes code for the expression of an FK-520 PKS that produces an FK-520 analog that lacks the C-19 to C-20 double bond of FK-520 and has a C-20 hydroxyl group. Such analogs are preferred neurotrophins, because they have little or no immunosuppressant activity. This recombinant fifth extender module coding sequence can be combined with other coding
20 sequences to make additional compounds of the invention. In an illustrative embodiment, the present invention provides a recombinant FK-520 PKS that contains both this fifth extender module and the recombinant fourth extender module described above that comprises the coding sequence for the fourth extender module AT domain of the FK-506 PKS. The invention also provides recombinant host cells derived from FK-506 producing
25 host cells that have been mutated to prevent production of FK-506 but that express this recombinant PKS and so synthesize the corresponding (lacking the C-19 to C-20 double bond of FK-506 and having a C-20 hydroxyl group) FK-506 derivative. In another embodiment, the present invention provides a recombinant FK-506 PKS in which the DH domain of module 5 has been deleted or otherwise rendered inactive and thus produces
30 this novel polyketide.

- 61 -

The sixth extender module of the FK-520 PKS includes a KS, an AT specific for methylmalonyl CoA, a KR, a DH, an ER, and an ACP. The recombinant DNA compounds of the invention that encode the sixth extender module of the FK-520 PKS and the corresponding polypeptides encoded thereby are useful for a variety of applications. In one embodiment, a DNA compound comprising a sequence that encodes the FK-520 sixth extender module is inserted into a DNA compound that comprises the coding sequence for a heterologous PKS. The resulting construct, in which the coding sequence for a module of the heterologous PKS is either replaced by that for the sixth extender module of the FK-520 PKS or the latter is merely added to coding sequences for the modules of the heterologous PKS, provides a novel PKS coding sequence. In another embodiment, a DNA compound comprising a sequence that encodes the sixth extender module of the FK-520 PKS is inserted into a DNA compound that comprises the coding sequence for the remainder of the FK-520 PKS or a recombinant FK-520 PKS that produces an FK-520 derivative.

In another embodiment, a portion of the sixth extender module coding sequence is utilized in conjunction with other PKS coding sequences to create a hybrid module. In this embodiment, the invention provides, for example, either replacing the methylmalonyl CoA specific AT with a malonyl CoA, ethylmalonyl CoA, or 2-hydroxymalonyl CoA specific AT; deleting any one, two, or all three of the KR, DH, and ER; and/or replacing any one, two, or all three of the KR, DH, and ER with another KR, DH, and ER. In addition, the KS and/or ACP can be replaced with another KS and/or ACP. In each of these replacements, the heterologous KS, AT, DH, KR, ER, or ACP coding sequence can originate from a coding sequence for another module of the FK-520 PKS, from a coding sequence for a PKS that produces a polyketide other than FK-520, or from chemical synthesis. The resulting heterologous sixth extender module coding sequence can be utilized in conjunction with a coding sequence for a PKS that synthesizes FK-520, an FK-520 derivative, or another polyketide. In similar fashion, the corresponding domains in a module of a heterologous PKS can be replaced by one or more domains of the sixth extender module of the FK-520 PKS.

- 62 -

In an illustrative embodiment, the present invention provides a set of recombinant FK-520 PKS genes in which the coding sequences for the DH and ER domains of the sixth extender module have been deleted or mutated to render them non-functional. In one such mutated gene, the KR, ER, and DH coding sequences are replaced with those encoding only a KR domain from another PKS gene. This can also be accomplished by simply replacing the coding sequences for extender module six with those for an extender module having a methylmalonyl specific AT and only a KR domain from a heterologous PKS gene, such as, for example, the coding sequences for extender module two encoded by the *eryAI* gene. The resulting PKS genes code for the expression of an FK-520 PKS that produces an FK-520 analog that has a C-18 hydroxyl group. Such analogs are preferred neurotrophins, because they have little or no immunosuppressant activity. This recombinant sixth extender module coding sequence can be combined with other coding sequences to make additional compounds of the invention. In an illustrative embodiment, the present invention provides a recombinant FK-520 PKS that contains both this sixth extender module and the recombinant fourth extender module described above that comprises the coding sequence for the fourth extender module AT domain of the FK-506 PKS. The invention also provides recombinant host cells derived from FK-506 producing host cells that have been mutated to prevent production of FK-506 but that express this recombinant PKS and so synthesize the corresponding (having a C-18 hydroxyl group) FK-506 derivative. In another embodiment, the present invention provides a recombinant FK-506 PKS in which the DH and ER domains of module 6 have been deleted or otherwise rendered inactive and thus produces this novel polyketide.

The seventh extender module of the FK-520 PKS includes a KS, an AT specific for 2-hydroxymalonyl CoA, a KR, a DH, an ER, and an ACP. The recombinant DNA compounds of the invention that encode the seventh extender module of the FK-520 PKS and the corresponding polypeptides encoded thereby are useful for a variety of applications. In one embodiment, a DNA compound comprising a sequence that encodes the FK-520 seventh extender module is inserted into a DNA compound that comprises the coding sequence for a heterologous PKS. The resulting construct, in which the coding sequence for a module of the heterologous PKS is either replaced by that for the seventh

- 63 -

extender module of the FK-520 PKS or the latter is merely added to coding sequences for the modules of the heterologous PKS, provides a novel PKS coding sequence. In another embodiment, a DNA compound comprising a sequence that encodes the seventh extender module of the FK-520 PKS is inserted into a DNA compound that comprises the coding
5 sequence for the remainder of the FK-520 PKS or a recombinant FK-520 PKS that produces an FK-520 derivative.

In another embodiment, a portion or all of the seventh extender module coding sequence is utilized in conjunction with other PKS coding sequences to create a hybrid module. In this embodiment, the invention provides, for example, either replacing the 2-
10 hydroxymalonyl CoA specific AT with a methylmalonyl CoA, ethylmalonyl CoA, or malonyl CoA specific AT; deleting the KR, the DH, and/or the ER; and/or replacing the KR, DH, and/or ER. In addition, the KS and/or ACP can be replaced with another KS and/or ACP. In each of these replacements or insertions, the heterologous KS, AT, DH, KR, ER, or ACP coding sequence can originate from a coding sequence for another
15 module of the FK-520 PKS, from a coding sequence for a PKS that produces a polyketide other than FK-520, or from chemical synthesis. The resulting heterologous seventh extender module coding sequence can be utilized in conjunction with a coding sequence for a PKS that synthesizes FK-520, an FK-520 derivative, or another polyketide. In similar fashion, the corresponding domains in a module of a heterologous PKS can be
20 replaced by one or more domains of the seventh extender module of the FK-520 PKS.

In an illustrative embodiment, the present invention provides a set of recombinant FK-520 PKS genes in which the coding sequences for the AT domain of the seventh extender module has been replaced with those encoding an AT domain for malonyl, methylmalonyl, or ethylmalonyl CoA from another PKS gene. The resulting PKS genes
25 code for the expression of an FK-520 PKS that produces an FK-520 analog that lacks the C-15 methoxy group, having instead a hydrogen, methyl, or ethyl group at that position, respectively. Such analogs are preferred, because they are more slowly metabolized than FK-520. This recombinant seventh extender module coding sequence can be combined with other coding sequences to make additional compounds of the invention. In an
30 illustrative embodiment, the present invention provides a recombinant FK-520 PKS that

- 64 -

contains both this seventh extender module and the recombinant fourth extender module described above that comprises the coding sequence for the fourth extender module AT domain of the FK-506 PKS. The invention also provides recombinant host cells derived from FK-506 producing host cells that have been mutated to prevent production of FK-506 but that express this recombinant PKS and so synthesize the corresponding (C-15-desmethoxy) FK-506 derivative. In another embodiment, the present invention provides a recombinant FK-506 PKS in which the AT domain of module 7 has been replaced and thus produces this novel polyketide.

In another illustrative embodiment, the present invention provides a hybrid PKS in which the AT and KR domains of module 7 of the FK-520 PKS are replaced by a methylmalonyl specific AT domain and an inactive KR domain, such as, for example, the AT and KR domains of extender module 6 of the rapamycin PKS. The resulting hybrid PKS produces 15-desmethoxy-15-methyl-16-oxo-FK-520, a neurotrophin compound.

The eighth extender module of the FK-520 PKS includes a KS, an AT specific for 2-hydroxymalonyl CoA, a KR, and an ACP. The recombinant DNA compounds of the invention that encode the eighth extender module of the FK-520 PKS and the corresponding polypeptides encoded thereby are useful for a variety of applications. In one embodiment, a DNA compound comprising a sequence that encodes the FK-520 eighth extender module is inserted into a DNA compound that comprises the coding sequence for a heterologous PKS. The resulting construct, in which the coding sequence for a module of the heterologous PKS is either replaced by that for the eighth extender module of the FK-520 PKS or the latter is merely added to coding sequences for the modules of the heterologous PKS, provides a novel PKS coding sequence. In another embodiment, a DNA compound comprising a sequence that encodes the eighth extender module of the FK-520 PKS is inserted into a DNA compound that comprises the coding sequence for the remainder of the FK-520 PKS or a recombinant FK-520 PKS that produces an FK-520 derivative.

In another embodiment, a portion of the eighth extender module coding sequence is utilized in conjunction with other PKS coding sequences to create a hybrid module. In this embodiment, the invention provides, for example, either replacing the 2-

- 65 -

hydroxymalonyl CoA specific AT with a methylmalonyl CoA, ethylmalonyl CoA, or malonyl CoA specific AT; deleting or replacing the KR; and/or inserting a DH or a DH and an ER. In addition, the KS and/or ACP can be replaced with another KS and/or ACP. In each of these replacements, the heterologous KS, AT, DH, KR, ER, or ACP coding
5 sequence can originate from a coding sequence for another module of the FK-520 PKS, from a coding sequence for a PKS that produces a polyketide other than FK-520, or from chemical synthesis. The resulting heterologous eighth extender module coding sequence can be utilized in conjunction with a PKS that synthesizes FK-520, an FK-520 derivative, or another polyketide. In similar fashion, the corresponding domains in a module of a
10 heterologous PKS can be replaced by one or more domains of the eighth extender module of the FK-520 PKS.

In an illustrative embodiment, the present invention provides a set of recombinant FK-520 PKS genes in which the coding sequences for the AT domain of the eighth extender module has been replaced with those encoding an AT domain for malonyl,
15 methylmalonyl, or ethylmalonyl CoA from another PKS gene. The resulting PKS genes code for the expression of an FK-520 PKS that produces an FK-520 analog that lacks the C-13 methoxy group, having instead a hydrogen, methyl, or ethyl group at that position, respectively. Such analogs are preferred, because they are more slowly metabolized than FK-520. This recombinant eighth extender module coding sequence can be combined
20 with other coding sequences to make additional compounds of the invention. In an illustrative embodiment, the present invention provides a recombinant FK-520 PKS that contains both this eighth extender module and the recombinant fourth extender module described above that comprises the coding sequence for the fourth extender module AT domain of the FK-506 PKS. The invention also provides recombinant host cells derived
25 from FK-506 producing host cells that have been mutated to prevent production of FK-506 but that express this recombinant PKS and so synthesize the corresponding (C-13-desmethoxy) FK-506 derivative. In another embodiment, the present invention provides a recombinant FK-506 PKS in which the AT domain of module 8 has been replaced and thus produces this novel polyketide.

- 66 -

The ninth extender module of the FK-520 PKS includes a KS, an AT specific for methylmalonyl CoA, a KR, a DH, an ER, and an ACP. The recombinant DNA compounds of the invention that encode the ninth extender module of the FK-520 PKS and the corresponding polypeptides encoded thereby are useful for a variety of applications. In one embodiment, a DNA compound comprising a sequence that encodes the FK-520 ninth extender module is inserted into a DNA compound that comprises the coding sequence for a heterologous PKS. The resulting construct, in which the coding sequence for a module of the heterologous PKS is either replaced by that for the ninth extender module of the FK-520 PKS or the latter is merely added to coding sequences for the modules of the heterologous PKS, provides a novel PKS coding sequence. In another embodiment, a DNA compound comprising a sequence that encodes the ninth extender module of the FK-520 PKS is inserted into a DNA compound that comprises the coding sequence for the remainder of the FK-520 PKS or a recombinant FK-520 PKS that produces an FK-520 derivative.

In another embodiment, a portion of the ninth extender module coding sequence is utilized in conjunction with other PKS coding sequences to create a hybrid module. In this embodiment, the invention provides, for example, either replacing the methylmalonyl CoA specific AT with a malonyl CoA, ethylmalonyl CoA, or 2-hydroxymalonyl CoA specific AT; deleting any one, two, or all three of the KR, DH, and ER; and/or replacing any one, two, or all three of the KR, DH, and ER with another KR, DH, and/or ER. In addition, the KS and/or ACP can be replaced with another KS and/or ACP. In each of these replacements, the heterologous KS, AT, DH, KR, ER, or ACP coding sequence can originate from a coding sequence for another module of the FK-520 PKS, from a coding sequence for a PKS that produces a polyketide other than FK-520, or from chemical synthesis. The resulting heterologous ninth extender module coding sequence can be utilized in conjunction with a PKS that synthesizes FK-520, an FK-520 derivative, or another polyketide. In similar fashion, the corresponding domains in a module of a heterologous PKS can be replaced by one or more domains of the ninth extender module of the FK-520 PKS.

- 67 -

The tenth extender module of the FK-520 PKS includes a KS, an AT specific for malonyl CoA, and an ACP. The recombinant DNA compounds of the invention that encode the tenth extender module of the FK-520 PKS and the corresponding polypeptides encoded thereby are useful for a variety of applications. In one embodiment, a DNA
5 compound comprising a sequence that encodes the FK-520 tenth extender module is inserted into a DNA compound that comprises the coding sequence for a heterologous PKS. The resulting construct, in which the coding sequence for a module of the heterologous PKS is either replaced by that for the tenth extender module of the FK-520 PKS or the latter is merely added to coding sequences for the modules of the
10 heterologous PKS, provides a novel PKS coding sequence. In another embodiment, a DNA compound comprising a sequence that encodes the tenth extender module of the FK-520 PKS is inserted into a DNA compound that comprises the coding sequence for the remainder of the FK-520 PKS or a recombinant FK-520 PKS that produces an FK-520 derivative.

15 In another embodiment, a portion or all of the tenth extender module coding sequence is utilized in conjunction with other PKS coding sequences to create a hybrid module. In this embodiment, the invention provides, for example, either replacing the malonyl CoA specific AT with a methylmalonyl CoA, ethylmalonyl CoA, or 2-hydroxymalonyl CoA specific AT; and/or inserting a KR, a KR and DH, or a KR, DH, and an ER. In addition, the KS and/or ACP can be replaced with another KS and/or ACP.
20 In each of these replacements or insertions, the heterologous KS, AT, DH, KR, ER, or ACP coding sequence can originate from a coding sequence for another module of the FK-520 PKS, from a coding sequence for a PKS that produces a polyketide other than FK-520, or from chemical synthesis. The resulting heterologous tenth extender module
25 coding sequence can be utilized in conjunction with a coding sequence for a PKS that synthesizes FK-520, an FK-520 derivative, or another polyketide. In similar fashion, the corresponding domains in a module of a heterologous PKS can be replaced by one or more domains of the tenth extender module of the FK-520 PKS.

30 The FK-520 polyketide precursor produced by the action of the tenth extender module of the PKS is then attached to pipecolic acid and cyclized to form FK-520. The

- 68 -

enzyme FkbP is the NRPS like enzyme that catalyzes these reactions. FkbP also includes a thioesterase activity that cleaves the nascent FK-520 polyketide from the NRPS. The present invention provides recombinant DNA compounds that encode the *fkbP* gene and so provides recombinant methods for expressing the *fkbP* gene product in recombinant host cells. The recombinant *fkbP* genes of the invention include those in which the coding sequence for the adenylation domain has been mutated or replaced with coding sequences from other NRPS like enzymes so that the resulting recombinant FkbP incorporates a moiety other than pipecolic acid. For the construction of host cells that do not naturally produce pipecolic acid, the present invention provides recombinant DNA compounds that express the enzymes that catalyze at least some of the biosynthesis of pipecolic acid (see Nielsen *et al.*, 1991, *Biochem. 30*: 5789-96). The *fkbL* gene encodes a homolog of RapL, a lysine cyclodeaminase responsible in part for producing the pipecolate unit added to the end of the polyketide chain. The *fkbB* and *fkbL* recombinant genes of the invention can be used in heterologous hosts to produce compounds such as FK-520 or, in conjunction with other PKS or NRPS genes, to produce known or novel polyketides and non-ribosomal peptides.

The present invention also provides recombinant DNA compounds that encode the P450 oxidase and methyltransferase genes involved in the biosynthesis of FK-520. Figure 2 shows the various sites on the FK-520 polyketide core structure at which these enzymes act. By providing these genes in recombinant form, the present invention provides recombinant host cells that can produce FK-520. This is accomplished by introducing the recombinant PKS, P450 oxidase, and methyltransferase genes into a heterologous host cell. In a preferred embodiment, the heterologous host cell is *Streptomyces coelicolor* CH999 or *Streptomyces lividans* K4-114, as described in U.S. Patent No. 5,830,750 and U.S. patent application Serial Nos. 08/828,898, filed 31 Mar. 1997, and 09/181,833, filed 28 Oct. 1998, each of which is incorporated herein by reference. In addition, by providing recombinant host cells that express only a subset of these genes, the present invention provides methods for making FK-520 precursor compounds not readily obtainable by other means.

- 69 -

In a related aspect, the present invention provides recombinant DNA compounds and vectors that are useful in generating, by homologous recombination, recombinant host cells that produce FK-520 precursor compounds. In this aspect of the invention, a native host cell that produces FK-520 is transformed with a vector (such as an SCP2*
5 derived vector for *Streptomyces* host cells) that encodes one or more disrupted genes (i.e., a hydroxylase, a methyltransferase, or both) or merely flanking regions from those genes. When the vector integrates by homologous recombination, the native, functional gene is deleted or replaced by the non-functional recombinant gene, and the resulting host cell thus produces an FK-520 precursor. Such host cells can also be complemented by
10 introduction of a modified form of the deleted or mutated non-functional gene to produce a novel compound.

In one important embodiment, the present invention provides a hybrid PKS and the corresponding recombinant DNA compounds that encode those hybrid PKS enzymes. For purposes of the present invention a hybrid PKS is a recombinant PKS that comprises
15 all or part of one or more modules and thioesterase/cyclase domain of a first PKS and all or part of one or more modules, loading module, and thioesterase/cyclase domain of a second PKS. In one preferred embodiment, the first PKS is all or part of the FK-520 PKS, and the second PKS is only a portion or all of a non-FK-520 PKS.

One example of the preferred embodiment is an FK-520 PKS in which the AT
20 domain of module 8, which specifies a hydroxymalonyl CoA and from which the C-13 methoxy group of FK-520 is derived, is replaced by an AT domain that specifies a malonyl, methylmalonyl, or ethylmalonyl CoA. Examples of such replacement AT domains include the AT domains from modules 3, 12, and 13 of the rapaymycin PKS and from modules 1 and 2 of the erythromycin PKS. Such replacements, conducted at the
25 level of the gene for the PKS, are illustrated in the examples below. Another illustrative example of such a hybrid PKS includes an FK-520 PKS in which the natural loading module has been replaced with a loading module of another PKS. Another example of such a hybrid PKS is an FK-520 PKS in which the AT domain of module three is replaced with an AT domain that binds methylmalonyl CoA.

- 70 -

In another preferred embodiment, the first PKS is most but not all of a non-FK-520 PKS, and the second PKS is only a portion or all of the FK-520 PKS. An illustrative example of such a hybrid PKS includes an erythromycin PKS in which an AT specific for methylmalonyl CoA is replaced with an AT from the FK-520 PKS specific for malonyl CoA.

Those of skill in the art will recognize that all or part of either the first or second PKS in a hybrid PKS of the invention need not be isolated from a naturally occurring source. For example, only a small portion of an AT domain determines its specificity. See U.S. provisional patent application Serial No. 60/091,526, incorporated herein by reference. The state of the art in DNA synthesis allows the artisan to construct *de novo* DNA compounds of size sufficient to construct a useful portion of a PKS module or domain. For purposes of the present invention, such synthetic DNA compounds are deemed to be a portion of a PKS.

Thus, the hybrid modules of the invention are incorporated into a PKS to provide a hybrid PKS of the invention. A hybrid PKS of the invention can result not only:

(i) from fusions of heterologous domain (where heterologous means the domains in that module are from at least two different naturally occurring modules) coding sequences to produce a hybrid module coding sequence contained in a PKS gene whose product is incorporated into a PKS,

but also:

(ii) from fusions of heterologous module (where heterologous module means two modules are adjacent to one another that are not adjacent to one another in naturally occurring PKS enzymes) coding sequences to produce a hybrid coding sequence contained in a PKS gene whose product is incorporated into a PKS,

(iii) from expression of one or more FK-520 PKS genes with one or more non-FK-520 PKS genes, including both naturally occurring and recombinant non-FK-520 PKS genes, and

(iv) from combinations of the foregoing.

Various hybrid PKSs of the invention illustrating these various alternatives are described herein.

- 71 -

Examples of the production of a hybrid PKS by co-expression of PKS genes from the FK-520 PKS and another non-FK-520 PKS include hybrid PKS enzymes produced by coexpression of FK-520 and rapamycin PKS genes. Preferably, such hybrid PKS enzymes are produced in recombinant *Streptomyces* host cells that produce FK-520 or FK-506 but have been mutated to inactivate the gene whose function is to be replaced by the rapamycin PKS gene introduced to produce the hybrid PKS. Particular examples include (i) replacement of the *fkbc* gene with the *rapB* gene; and (ii) replacement of the *fkba* gene with the *rapC* gene. The latter hybrid PKS produces 13,15-didesmethoxy-FK-520, if the host cell is an FK-520 producing host cell, and 13,15-didesmethoxy-FK-506, if the host cell is an FK-506 producing host cell. The compounds produced by these hybrid PKS enzymes are immunosuppressants and neurotrophins but can be readily modified to act only as neurotrophins, as described in Example 6, below.

Other illustrative hybrid PKS enzymes of the invention are prepared by replacing the *fkba* gene of an FK-520 or FK-506 producing host cell with a hybrid *fkba* gene in which: (a) the extender module 8 through 10, inclusive, coding sequences have been replaced by the coding sequences for extender modules 12 to 14, inclusive, of the rapamycin PKS; and (b) the module 8 coding sequences have been replaced by the module 8 coding sequence of the rifamycin PKS. When expressed with the other, naturally occurring FK-520 or FK-506 PKS genes and the genes of the modification enzymes, the resulting hybrid PKS enzymes produce, respectively, (a) 13-desmethoxy-FK-520 or 13-desmethoxy-FK-506; and (b) 13-desmethoxy-13-methyl-FK-520 or 13-desmethoxy-13-methyl-FK-506. In a preferred embodiment, these recombinant PKS genes of the invention are introduced into the producing host cell by a vector such as pHU204, which is a plasmid pRM5 derivative that has the well-characterized SCP2* replicon, the *colE1* replicon, the *tsr* and *bla* resistance genes, and a *cos* site. This vector can be used to introduce the recombinant *fkba* replacement gene in an FK-520 or FK-506 producing host cell (or a host cell derived therefrom in which the endogenous *fkba* gene has either been rendered inactive by mutation, deletion or homologous recombination with the gene that replaces it) to produce the desired hybrid PKS.

- 72 -

In constructing hybrid PKSs of the invention, certain general methods may be helpful. For example, it is often beneficial to retain the framework of the module to be altered to make the hybrid PKS. Thus, if one desires to add DH and ER functionalities to a module, it is often preferred to replace the KR domain of the original module with a KR, DH, and ER domain-containing segment from another module, instead of merely inserting DH and ER domains. One can alter the stereochemical specificity of a module by replacement of the KS domain with a KS domain from a module that specifies a different stereochemistry. See Lau *et al.*, 1999, "Dissecting the role of acyltransferase domains of modular polyketide synthases in the choice and stereochemical fate of extender units," *Biochemistry* 38(5):1643-1651, incorporated herein by reference. Stereochemistry can also be changed by changing the KR domain. Also, one can alter the specificity of an AT domain by changing only a small segment of the domain. See Lau *et al.*, *supra*. One can also take advantage of known linker regions in PKS proteins to link modules from two different PKSs to create a hybrid PKS. See Gokhale *et al.*, 16 Apr. 1999, "Dissecting and Exploiting Intermodular Communication in Polyketide Synthases," *Science* 284: 482-485, incorporated herein by reference.

The following Table lists references describing illustrative PKS genes and corresponding enzymes that can be utilized in the construction of the recombinant PKSs and the corresponding DNA compounds that encode them of the invention. Also presented are various references describing tailoring enzymes and corresponding genes that can be employed in accordance with the methods of the present invention.

Avermectin

U.S. Pat. No. 5,252,474 to Merck.

MacNeil *et al.*, 1993, Industrial Microorganisms: Basic and Applied Molecular Genetics, Baltz, Hegeman, & Skatrud, eds. (ASM), pp. 245-256, A Comparison of the Genes Encoding the Polyketide Synthases for Avermectin, Erythromycin, and Nemadectin.

MacNeil *et al.*, 1992, *Gene* 115: 119-125, Complex Organization of the *Streptomyces avermitilis* genes encoding the avermectin polyketide synthase.

- 73 -

Ikeda *et al.*, Aug. 1999, Organization of the biosynthetic gene cluster for the polyketide anthelmintic macrolide avermectin in *Streptomyces avermitilis*, *Proc. Natl. Acad. Sci. USA* 96: 9509-9514.

Candicidin (FR008)

5 Hu *et al.*, 1994, *Mol. Microbiol.* 14: 163-172.

Epothilone

U.S. Pat. App. Serial No. 60/130,560, filed 22 April 1999.

Erythromycin

PCT Pub. No. 93/13663 to Abbott.

10 US Pat. No. 5,824,513 to Abbott.

Donadio *et al.*, 1991, *Science* 252:675-9.

Cortes *et al.*, 8 Nov. 1990, *Nature* 348:176-8, An unusually large multifunctional polypeptide in the erythromycin producing polyketide synthase of *Saccharopolyspora erythraea*.

15 Glycosylation Enzymes

PCT Pat. App. Pub. No. 97/23630 to Abbott.

FK-506

Motamedi *et al.*, 1998, The biosynthetic gene cluster for the macrolactone ring of the immunosuppressant FK-506, *Eur. J. biochem.* 256: 528-534.

20 Motamedi *et al.*, 1997, Structural organization of a multifunctional polyketide synthase involved in the biosynthesis of the macrolide immunosuppressant FK-506, *Eur. J. Biochem.* 244: 74-80.

Methyltransferase

25 US 5,264,355, issued 23 Nov. 1993, Methylating enzyme from *Streptomyces* MA6858. 31-O-desmethyl-FK-506 methyltransferase.

Motamedi *et al.*, 1996, Characterization of methyltransferase and hydroxylase genes involved in the biosynthesis of the immunosuppressants FK-506 and FK-520, *J. Bacteriol.* 178: 5243-5248.

Streptomyces hygroscopicus

30 U.S. patent application Serial No. 09/154,083, filed 16 Sep. 1998.

- 74 -

Lovastatin

U.S. Pat. No. 5,744,350 to Merck.

Narbomycin

U.S. patent application Serial No. 60/107,093, filed 5 Nov. 1998, and Serial No.
5 60/120,254, filed 16 Feb. 1999.

Nemadectin

MacNeil *et al.*, 1993, *supra*.

Niddamycin

Kakavas *et al.*, 1997, Identification and characterization of the niddamycin
10 polyketide synthase genes from *Streptomyces caelestis*, *J. Bacteriol.* 179: 7515-7522.

Oleandomycin

Swan *et al.*, 1994, Characterisation of a *Streptomyces antibioticus* gene encoding
a type I polyketide synthase which has an unusual coding sequence, *Mol. Gen. Genet.*
242: 358-362.

15 U.S. patent application Serial No. 60/120,254, filed 16 Feb. 1999.

Olano *et al.*, 1998, Analysis of a *Streptomyces antibioticus* chromosomal region
involved in oleandomycin biosynthesis, which encodes two glycosyltransferases
responsible for glycosylation of the macrolactone ring, *Mol. Gen. Genet.* 259(3): 299-
308.

20 **Picromycin**

PCT patent application US99/15047, filed 2 Jul. 1999.

Xue *et al.*, 1998, Hydroxylation of macrolactones YC-17 and narbomycin is
mediated by the *pikC*-encoded cytochrome P450 in *Streptomyces venezuelae*, *Chemistry*
& *Biology* 5(11): 661-667.

25 Xue *et al.*, Oct. 1998, A gene cluster for macrolide antibiotic biosynthesis in
Streptomyces venezuelae: Architecture of metabolic diversity, *Proc. Natl. Acad. Sci.*
USA 95: 12111 12116.

Platenolide

EP Pat. App. Pub. No. 791,656 to Lilly.

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- 75 -

Rapamycin

Schwecke *et al.*, Aug. 1995, The biosynthetic gene cluster for the polyketide rapamycin, *Proc. Natl. Acad. Sci. USA* 92:7839-7843.

Aparicio *et al.*, 1996, Organization of the biosynthetic gene cluster for rapamycin
5 in *Streptomyces hygroscopicus*: analysis of the enzymatic domains in the modular polyketide synthase, *Gene* 169: 9-16.

Rifamycin

August *et al.*, 13 Feb. 1998, Biosynthesis of the ansamycin antibiotic rifamycin: deductions from the molecular analysis of the *rif* biosynthetic gene cluster of
10 *Amycolatopsis mediterranei* S669, *Chemistry & Biology*, 5(2): 69-79.

Sorangium PKS

U.S. patent application Serial No. 09/144,085, filed 31 Aug. 1998.

Soraphen

U.S. Pat. No. 5,716,849 to Novartis.

15 Schupp *et al.*, 1995, *J. Bacteriology* 177: 3673-3679. A *Sorangium cellulosum* (Myxobacterium) Gene Cluster for the Biosynthesis of the Macrolide Antibiotic Soraphen A: Cloning, Characterization, and Homology to Polyketide Synthase Genes from Actinomycetes.

Spiramycin

20 U.S. Pat. No. 5,098,837 to Lilly.

Activator Gene

U.S. Pat. No. 5,514,544 to Lilly.

Tylosin

EP Pub. No. 791,655 to Lilly.

25 U.S. Pat. No. 5,876,991 to Lilly.

Kuhstoss *et al.*, 1996, *Gene* 183:231-6., Production of a novel polyketide through the construction of a hybrid polyketide synthase.

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- 76 -

Tailoring enzymes

Merson-Davies and Cundliffe, 1994, *Mol. Microbiol.* 13: 349-355. Analysis of five tylosin biosynthetic genes from the *tylBA* region of the *Streptomyces fradiae* genome.

5 As the above Table illustrates, there are a wide variety of polyketide synthase genes that serve as readily available sources of DNA and sequence information for use in constructing the hybrid PKS-encoding DNA compounds of the invention. Methods for constructing hybrid PKS-encoding DNA compounds are described without reference to the FK-520 PKS in PCT patent publication No. 98/51695; U.S. Patent Nos. 5,672,491
10 and 5,712,146 and U.S. patent application Serial Nos. 09/073,538, filed 6 May 1998, and 09/141,908, filed 28 Aug 1998, each of which is incorporated herein by reference.

The hybrid PKS-encoding DNA compounds of the invention can be and often are hybrids of more than two PKS genes. Moreover, there are often two or more modules in the hybrid PKS in which all or part of the module is derived from a second (or third)
15 PKS. Thus, as one illustrative example, the present invention provides a hybrid FK-520 PKS that contains the naturally occurring loading module and FkbP as well as modules one, two, four, six, seven, and eight, nine, and ten of the FK-520 PKS and further contains hybrid or heterologous modules three and five. Hybrid or heterologous module three contains an AT domain that is specific of methylmalonyl CoA and can be derived
20 for example, from the erythromycin or rapamycin PKS genes. Hybrid or heterologous module five contains an AT domain that is specific for malonyl CoA and can be derived for example, from the picromycin or rapamycin PKS genes.

While an important embodiment of the present invention relates to hybrid PKS enzymes and corresponding genes, the present invention also provides recombinant FK-
25 520 PKS genes in which there is no second PKS gene sequence present but which differ from the FK-520 PKS gene by one or more deletions. The deletions can encompass one or more modules and/or can be limited to a partial deletion within one or more modules. When a deletion encompasses an entire module, the resulting FK-520 derivative is at least two carbons shorter than the gene from which it was derived. When a deletion is
30 within a module, the deletion typically encompasses a KR, DH, or ER domain, or both

- 77 -

DH and ER domains, or both KR and DH domains, or all three KR, DH, and ER domains.

To construct a hybrid PKS or FK-520 derivative PKS gene of the invention, one can employ a technique, described in PCT Pub. No. 98/27203 and U.S. patent application
5 Serial No. 08/989,332, filed 11 Dec. 1997, each of which is incorporated herein by reference, in which the large PKS gene is divided into two or more, typically three, segments, and each segment is placed on a separate expression vector. In this manner, each of the segments of the gene can be altered, and various altered segments can be combined in a single host cell to provide a recombinant PKS gene of the invention. This
10 technique makes more efficient the construction of large libraries of recombinant PKS genes, vectors for expressing those genes, and host cells comprising those vectors.

Thus, in one important embodiment, the recombinant DNA compounds of the invention are expression vectors. As used herein, the term expression vector refers to any nucleic acid that can be introduced into a host cell or cell-free transcription and
15 translation medium. An expression vector can be maintained stably or transiently in a cell, whether as part of the chromosomal or other DNA in the cell or in any cellular compartment, such as a replicating vector in the cytoplasm. An expression vector also comprises a gene that serves to produce RNA that is translated into a polypeptide in the cell or cell extract. Furthermore, expression vectors typically contain additional
20 functional elements, such as resistance-conferring genes to act as selectable markers.

The various components of an expression vector can vary widely, depending on the intended use of the vector. In particular, the components depend on the host cell(s) in which the vector will be used or is intended to function. Vector components for expression and maintenance of vectors in *E. coli* are widely known and commercially
25 available, as are vector components for other commonly used organisms, such as yeast cells and *Streptomyces* cells.

In a preferred embodiment, the expression vectors of the invention are used to construct recombinant *Streptomyces* host cells that express a recombinant PKS of the invention. Preferred *Streptomyces* host cell/vector combinations of the invention include
30 *S. coelicolor* CH999 and *S. lividans* K4-114 host cells, which do not produce

- 78 -

actinorhodin, and expression vectors derived from the pRM1 and pRM5 vectors, as described in U.S. Patent No. 5,830,750 and U.S. patent application Serial Nos. 08/828,898, filed 31 Mar. 1997, and 09/181,833, filed 28 Oct. 1998, each of which is incorporated herein by reference.

5 The present invention provides a wide variety of expression vectors for use in *Streptomyces*. For replicating vectors, the origin of replication can be, for example and without limitation, a low copy number vector, such as SCP2* (see Hopwood *et al.*, *Genetic Manipulation of Streptomyces: A Laboratory manual* (The John Innes Foundation, Norwich, U.K., 1985); Lydiate *et al.*, 1985, *Gene* 35: 223-235; and Kieser and Melton, 1988, *Gene* 65: 83-91, each of which is incorporated herein by reference),
10 SLP1.2 (Thompson *et al.*, 1982, *Gene* 20: 51-62, incorporated herein by reference), and SG5(ts) (Muth *et al.*, 1989, *Mol. Gen. Genet.* 219: 341-348, and Bierman *et al.*, 1992, *Gene* 116: 43-49, each of which is incorporated herein by reference), or a high copy number vector, such as pIJ101 and pJV1 (see Katz *et al.*, 1983, *J. Gen. Microbiol.* 129: 2703-2714; Vara *et al.*, 1989, *J. Bacteriol.* 171: 5782-5781; and Servin-Gonzalez, 1993, *Plasmid* 30: 131-140, each of which is incorporated herein by reference). Generally,
15 however, high copy number vectors are not preferred for expression of genes contained on large segments of DNA. For non-replicating and integrating vectors, it is useful to include at least an *E. coli* origin of replication, such as from pUC, pIP, pII, and pBR. For
20 phage based vectors, the phages phiC31 and KC515 can be employed (see Hopwood *et al.*, *supra*).

Typically, the expression vector will comprise one or more marker genes by which host cells containing the vector can be identified and/or selected. Useful antibiotic resistance conferring genes for use in *Streptomyces* host cells include the *ermE* (confers
25 resistance to erythromycin and other macrolides and lincomycin), *tsr* (confers resistance to thiostrepton), *aadA* (confers resistance to spectinomycin and streptomycin), *aacC4* (confers resistance to apramycin, kanamycin, gentamicin, geneticin (G418), and neomycin), *hyg* (confers resistance to hygromycin), and *vph* (confers resistance to viomycin) resistance conferring genes.

- 79 -

The recombinant PKS gene on the vector will be under the control of a promoter, typically with an attendant ribosome binding site sequence. The present invention provides the endogenous promoters of the FK-520 PKS and related biosynthetic genes in recombinant form, and these promoters are preferred for use in the native hosts and in
5 heterologous hosts in which the promoters function. A preferred promoter of the invention is the *fkfO* gene promoter, comprised in a sequence of about 270 bp between the start of the open reading frames of the *fkfO* and *fkfB* genes. The *fkfO* promoter is believed to be bi-directional in that it promotes transcription of the genes *fkfO*, *fkfP*, and *fkfA* in one direction and *fkfB*, *fkfC*, and *fkfL* in the other. Thus, in one aspect, the
10 present invention provides a recombinant expression vector comprising the promoter of the *fkfO* gene of an FK-520 producing organism positioned to transcribe a gene other than *fkfO*. In a preferred embodiment the transcribed gene is an FK-520 PKS gene. In another preferred embodiment, the transcribed gene is a gene that encodes a protein comprised in a hybrid PKS.

15 Heterologous promoters can also be employed and are preferred for use in host cells in which the endogenous FK-520 PKS gene promoters do not function or function poorly. A preferred heterologous promoter is the *actI* promoter and its attendant activator gene *actII-ORF4*, which is provided in the pRM1 and pRM5 expression vectors, *supra*. This promoter is activated in the stationary phase of growth when secondary metabolites
20 are normally synthesized. Other useful *Streptomyces* promoters include without limitation those from the *ermE* gene and the *melC1* gene, which act constitutively, and the *tipA* gene and the *merA* gene, which can be induced at any growth stage. In addition, the T7 RNA polymerase system has been transferred to *Streptomyces* and can be employed in the vectors and host cells of the invention. In this system, the coding sequence for the T7
25 RNA polymerase is inserted into a neutral site of the chromosome or in a vector under the control of the inducible *merA* promoter, and the gene of interest is placed under the control of the T7 promoter. As noted above, one or more activator genes can also be employed to enhance the activity of a promoter. Activator genes in addition to the *actII-ORF4* gene discussed above include *dnrI*, *redD*, and *ptpA* genes (see U.S. patent
30 application Serial No. 09/181,833, *supra*) to activate promoters under their control.

- 80 -

In addition to providing recombinant DNA compounds that encode the FK-520 PKS, the present invention also provides DNA compounds that encode the ethylmalonyl CoA and 2-hydroxymalonyl CoA utilized in the synthesis of FK-520. Thus, the present invention also provides recombinant host cells that express the genes required for the biosynthesis of ethylmalonyl CoA and 2-hydroxymalonyl CoA. Figures 3 and 4 show the location of these genes on the cosmids of the invention and the biosynthetic pathway that produces ethylmalonyl CoA.

For 2-hydroxymalonyl CoA biosynthesis, the *fkfH*, *fkfI*, *fkfJ*, and *fkfK* genes are sufficient to confer this ability on *Streptomyces* host cells. For conversion of 2-hydroxymalonyl to 2-methoxymalonyl, the *fkfG* gene is also employed. While the complete coding sequence for *fkfH* is provided on the cosmids of the invention, the sequence for this gene provided herein may be missing a T residue, based on a comparison made with a similar gene cloned from the ansamitocin gene cluster by Dr. H. Floss. Where the sequence herein shows one T, there may be two, resulting in an extension of the *fkfH* reading frame to encode the amino acid sequence:

MTIVKCLVWDLNLTWRGTVLEDDEVVLTDEIREVITTLDDRGILQAVASKNDH
DLAWERLERLGVAEYFVLARIGWGPKSQSVREIATELNFAPTTIAFIDDQPAERA
EVAFHLPEVRCYPAEQAATLLSLPEFSPPVSTVDSRRRLMYQAGFARDQAREA
YSGPDEDFLRSLDSMTIAPAGEEELSRVEELTLRTSQMNATGVHYSDADLRALL
TDPAHEVLVVTMGDRFGPHGAVGILLEKKPSTWHLKLLATSCRVV SFGAGATIL
NWLTDQGARAGAHLVADFRRTDRNRMM EIA YRFAGFADSDCPCVSEVAGASA
AGVERLHLEPSARPAPTTLTLTAADIAPVTVSAAG.

For ethylmalonyl CoA biosynthesis, one requires only a crotonyl CoA reductase, which can be supplied by the host cell but can also be supplied by recombinant expression of the *fkfS* gene of the present invention. To increase yield of ethylmalonyl CoA, one can also express the *fkfE* and *fkfU* genes as well. While such production can be achieved using only the recombinant genes above, one can also achieve such production by placing into the recombinant host cell a large segment of the DNA provided by the cosmids of the invention. Thus, for 2-hydroxymalonyl and 2-methoxymalonyl CoA biosynthesis, one can simply provide the cells with the segment of

- 81 -

DNA located on the left side of the FK-520 PKS genes shown in Figure 1. For ethylmalonyl CoA biosynthesis, one can simply provide the cells with the segment of DNA located on the right side of the FK-520 PKS genes shown in Figure 1 or, alternatively, both the right and left segments of DNA.

5 The recombinant DNA expression vectors that encode these genes can be used to construct recombinant host cells that can make these important polyketide building blocks from cells that otherwise are unable to produce them. For example, *Streptomyces coelicolor* and *Streptomyces lividans* do not synthesize ethylmalonyl CoA or 2-hydroxymalonyl CoA. The invention provides methods and vectors for constructing
10 recombinant *Streptomyces coelicolor* and *Streptomyces lividans* that are able to synthesize either or both ethylmalonyl CoA and 2-hydroxymalonyl CoA. These host cells are thus able to make polyketides, those requiring these substrates, that cannot otherwise be made in such cells.

 In a preferred embodiment, the present invention provides recombinant
15 *Streptomyces* host cells, such as *S. coelicolor* and *S. lividans*, that have been transformed with a recombinant vector of the invention that codes for the expression of the ethylmalonyl CoA biosynthetic genes. The resulting host cells produce ethylmalonyl CoA and so are preferred host cells for the production of polyketides produced by PKS enzymes that comprise one or more AT domains specific for ethylmalonyl CoA.
20 Illustrative PKS enzymes of this type include the FK-520 PKS and a recombinant PKS in which one or more AT domains is specific for ethylmalonyl CoA.

 In a related embodiment, the present invention provides *Streptomyces* host cells in which one or more of the ethylmalonyl or 2-hydroxymalonyl biosynthetic genes have been deleted by homologous recombination or rendered inactive by mutation. For
25 example, deletion or inactivation of the *fkfG* gene can prevent formation of the methoxyl groups at C-13 and C-15 of FK-520 (or, in the corresponding FK-506 producing cell, FK-506), leading to the production of 13,15-didesmethoxy-13,15-dihydroxy-FK-520 (or, in the corresponding FK-506 producing cell, 13,15-didesmethoxy-13,15-dihydroxy-FK-506). If the *fkfG* gene product acts on 2-hydroxymalonyl and the resulting 2-
30 methoxymalonyl substrate is required for incorporation by the PKS, the AT domains of

- 82 -

modules 7 and 8 may bind malonyl CoA and methylmalonyl CoA. Such incorporation results in the production of a mixture of polyketides in which the methoxy groups at C-13 and C-15 of FK-520 (or FK-506) are replaced by either hydrogen or methyl.

5 This possibility of non-specific binding results from the construction of a hybrid PKS of the invention in which the AT domain of module 8 of the FK-520 PKS replaced the AT domain of module 6 of DEBS. The resulting PKS produced, in *Streptomyces lividans*, 6-dEB and 2-desmethyl-6-dEB, indicating that the AT domain of module 8 of the FK-520 PKS could bind malonyl CoA and methylmalonyl CoA substrates. Thus, one could possibly also prepare the 13,15-didesmethoxy-FK-520 and corresponding FK-506
10 compounds of the invention by deleting or otherwise inactivating one or more or all of the genes required for 2-hydroxymalonyl CoA biosynthesis, i.e., the *fkfH*, *fkfI*, *fkfJ*, and *fkfK* genes. In any event, the deletion or inactivation of one or more biosynthetic genes required for ethylmalonyl and/or 2-hydroxymalonyl production prevents the formation of polyketides requiring ethylmalonyl and/or 2-hydroxymalonyl for biosynthesis, and the
15 resulting host cells are thus preferred for production of polyketides that do not require the same.

The host cells of the invention can be grown and fermented under conditions known in the art for other purposes to produce the compounds of the invention. See, e.g., U.S. Patent Nos. 5,194,378; 5,116,756; and 5,494,820, incorporated herein by reference,
20 for suitable fermentation processes. The compounds of the invention can be isolated from the fermentation broths of these cultured cells and purified by standard procedures. Preferred compounds of the invention include the following compounds: 13-desmethoxy-FK-506; 13-desmethoxy-FK-520; 13,15-didesmethoxy-FK-506; 13,15-didesmethoxy-FK-520; 13-desmethoxy-18-hydroxy-FK-506; 13-desmethoxy-18-hydroxy-FK-520;
25 13,15-didesmethoxy-18-hydroxy-FK-506; and 13,15-didesmethoxy-18-hydroxy-FK-520. These compounds can be further modified as described for tacrolimus and FK-520 in U.S. Patent Nos. 5,225,403; 5,189,042; 5,164,495; 5,068,323; 4,980,466; and 4,920,218, incorporated herein by reference.

Other compounds of the invention are shown in Figure 8, Parts A and B. In Figure
30 8, Part A, illustrative C-32-substituted compounds of the invention are shown in two

- 83 -

columns under the heading R. The substituted compounds are preferred for topical administration and are applied to the dermis for treatment of conditions such as psoriasis. In Figure 8, Part B, illustrative reaction schemes for making the compounds shown in Figure 8, Part A, are provided. In the upper scheme in Figure 8, Part B, the C-32
5 substitution is a tetrazole moiety, illustrative of the groups shown in the left column under R in Figure 8, Part A. In the lower scheme in Figure 8, Part B, the C-32 substitution is a disubstituted amino group, where R₃ and R₄ can be any group similar to the illustrative groups shown attached to the amine in the right column under R in Figure 8, Part A. While Figure 8 shows the C-32-substituted compounds in which the C-15-methoxy is present, the invention includes these C-32-substituted compounds in which C-
10 15 is ethyl, methyl, or hydrogen. Also, while C-21 is shown as substituted with ethyl or allyl, the compounds of the invention includes the C-32-substituted compounds in which C-21 is substituted with hydrogen or methyl.

To make these C-32-substituted compounds, Figure 8, Part B, provides illustrative
15 reaction schemes. Thus, a selective reaction of the starting compound (see Figure 8, Part B, for an illustrative starting compound) with trifluoromethanesulfonic anhydride in the presence of a base yields the C-32 O-triflate derivative, as shown in the upper scheme of Figure 8, Part B. Displacement of the triflate with 1H-tetrazole or triazole derivatives provides the C-32 tetrazole or teiazole derivative. As shown in the lower scheme of
20 Figure 8, Part B, reacting the starting compound with p-nitrophenylchloroformate yields the corresponding carbonate, which, upon displacement with an amino compound, provides the corresponding carbamate derivative.

The compounds can be readily formulated to provide the pharmaceutical compositions of the invention. The pharmaceutical compositions of the invention can be
25 used in the form of a pharmaceutical preparation, for example, in solid, semisolid, or liquid form. This preparation contains one or more of the compounds of the invention as an active ingredient in admixture with an organic or inorganic carrier or excipient suitable for external, enteral, or parenteral application. The active ingredient may be compounded, for example, with the usual non-toxic, pharmaceutically acceptable carriers
30 for tablets, pellets, capsules, suppositories, solutions, emulsions, suspensions, and any

- 84 -

other form suitable for use. Suitable formulation processes and compositions for the compounds of the present invention are described with respect to tacrolimus in U.S. Patent Nos. 5,939,427; 5,922,729; 5,385,907; 5,338,684; and 5,260,301, incorporated herein by reference. Many of the compounds of the invention contain one or more chiral
5 centers, and all of the stereoisomers are included within the scope of the invention, as pure compounds as well as mixtures of stereoisomers. Thus the compounds of the invention may be supplied as a mixture of stereoisomers in any proportion.

The carriers which can be used include water, glucose, lactose, gum acacia, gelatin, mannitol, starch paste, magnesium trisilicate, talc, corn starch, keratin, colloidal
10 silica, potato starch, urea, and other carriers suitable for use in manufacturing preparations, in solid, semi-solid, or liquified form. In addition, auxiliary stabilizing, thickening, and coloring agents and perfumes may be used. For example, the compounds of the invention may be utilized with hydroxypropyl methylcellulose essentially as described in U.S. Patent No. 4,916,138, incorporated herein by reference, or with a
15 surfactant essentially as described in EPO patent publication No. 428,169, incorporated herein by reference.

Oral dosage forms may be prepared essentially as described by Hondo *et al.*, 1987, *Transplantation Proceedings XIX*, Supp. 6: 17-22, incorporated herein by reference. Dosage forms for external application may be prepared essentially as described
20 in EPO patent publication No. 423,714, incorporated herein by reference. The active compound is included in the pharmaceutical composition in an amount sufficient to produce the desired effect upon the disease process or condition.

For the treatment of conditions and diseases relating to immunosuppression or neuronal damage, a compound of the invention may be administered orally, topically,
25 parenterally, by inhalation spray, or rectally in dosage unit formulations containing conventional non-toxic pharmaceutically acceptable carriers, adjuvant, and vehicles. The term parenteral, as used herein, includes subcutaneous injections, and intravenous, intramuscular, and intrasternal injection or infusion techniques.

Dosage levels of the compounds of the present invention are of the order from
30 about 0.01 mg to about 50 mg per kilogram of body weight per day, preferably from

- 85 -

about 0.1 mg to about 10 mg per kilogram of body weight per day. The dosage levels are useful in the treatment of the above-indicated conditions (from about 0.7 mg to about 3.5 mg per patient per day, assuming a 70 kg patient). In addition, the compounds of the present invention may be administered on an intermittent basis, i.e., at semi-weekly,
5 weekly, semi-monthly, or monthly intervals.

The amount of active ingredient that may be combined with the carrier materials to produce a single dosage form will vary depending upon the host treated and the particular mode of administration. For example, a formulation intended for oral administration to humans may contain from 0.5 mg to 5 g of active agent compounded
10 with an appropriate and convenient amount of carrier material, which may vary from about 5 percent to about 95 percent of the total composition. Dosage unit forms will generally contain from about 0.5 mg to about 500 mg of active ingredient. For external administration, the compounds of the invention can be formulated within the range of, for example, 0.00001% to 60% by weight, preferably from 0.001% to 10% by weight, and
15 most preferably from about 0.005% to 0.8% by weight. The compounds and compositions of the invention are useful in treating disease conditions using doses and administration schedules as described for tacrolimus in U.S. Patent Nos. 5,542,436; 5,365,948; 5,348,966; and 5,196,437, incorporated herein by reference. The compounds of the invention can be used as single therapeutic agents or in combination with other
20 therapeutic agents. Drugs that can be usefully combined with compounds of the invention include one or more immunosuppressant agents such as rapamycin, cyclosporin A, FK-506, or one or more neurotrophic agents.

It will be understood, however, that the specific dosage level for any particular patient will depend on a variety of factors. These factors include the activity of the
25 specific compound employed; the age, body weight, general health, sex, and diet of the subject; the time and route of administration and the rate of excretion of the drug; whether a drug combination is employed in the treatment; and the severity of the particular disease or condition for which therapy is sought.

- 86 -

A detailed description of the invention having been provided above, the following examples are given for the purpose of illustrating the present invention and shall not be construed as being a limitation on the scope of the invention or claims.

5

Example 1

Replacement of Methoxyl with Hydrogen or Methyl at C-13 of FK-520

The C-13 methoxyl group is introduced into FK-520 via an AT domain in extender module 8 of the PKS that is specific for hydroxymalonyl and by methylation of the hydroxyl group by an S-adenosyl methionine (SAM) dependent methyltransferase.

10 Metabolism of FK-506 and FK-520 primarily involves oxidation at the C-13 position into an inactive derivative that is further degraded by host P450 and other enzymes. The present invention provides compounds related in structure to FK-506 and FK-520 that do not contain the C-13 methoxy group and exhibit greater stability and a longer half-life *in vivo*. These compounds are useful medicaments due to their immunosuppressive and
15 neurotrophic activities, and the invention provides the compounds in purified form and as pharmaceutical compositions.

The present invention also provides the novel PKS enzymes that produce these novel compounds as well as the expression vectors and host cells that produce the novel PKS enzymes. The novel PKS enzymes include, among others, those that contain an AT
20 domain specific for either malonyl CoA or methylmalonyl CoA in module 8 of the FK-506 and FK-520 PKS. This example describes the construction of recombinant DNA compounds that encode the novel FK-520 PKS enzymes and the transformation of host cells with those recombinant DNA compounds to produce the novel PKS enzymes and the polyketides produced thereby.

25 To construct an expression cassette for performing module 8 AT domain replacements in the FK-520 PKS, a 4.6 kb *Sph*I fragment from the FK-520 gene cluster was cloned into plasmid pLitmus 38 (a cloning vector available from New England Biolabs). The 4.6 kb *Sph*I fragment, which encodes the ACP domain of module 7 followed by module 8 through the KR domain, was isolated from an agarose gel after
30 digesting the cosmid pKOS65-C31 with *Sph* I. The clone having the insert oriented so

- 87 -

the single *SacI* site was nearest to the *SpeI* end of the polylinker was identified and designated as plasmid pKOS60-21-67. To generate appropriate cloning sites, two linkers were ligated sequentially as follows. First, a linker was ligated between the *SpeI* and *SacI* sites to introduce a *BglII* site at the 5' end of the cassette, to eliminate interfering polylinker sites, and to reduce the total insert size to 4.5 kb (the limit of the phage KC515). The ligation reactions contained 5 picomolar unphosphorylated linker DNA and 0.1 picomolar vector DNA, i.e., a 50-fold molar excess of linker to vector. The linker had the following sequence:

5'-CTAGTGGGCAGATCTGGCAGCT-3'
3'-ACCCGTCTAGACCG-5'

The resulting plasmid was designated pKOS60-27-1.

Next, a linker of the following sequence was ligated between the unique *SphI* and *AflIII* sites of plasmid pKOS60-27-1 to introduce an *NsiI* site at the 3' end of the module 8 cassette. The linker employed was:

5'-GGGATGCATGGC-3'
3'-GTACCCCTACGTACCGAATT-5'

The resulting plasmid was designated pKOS60-29-55.

To allow in-frame insertions of alternative AT domains, sites were engineered at the 5' end (*Avr II* or *Nhe I*) and 3' end (*Xho I*) of the AT domain using the polymerase chain reaction (PCR) as follows. Plasmid pKOS60-29-55 was used as a template for the PCR and sequence 5' to the AT domain was amplified with the primers *SpeBgl*-fwd and either *Avr*-rev or *Nhe*-rev:

SpeBgl-fwd 5'-CGACTCACTAGTGGGCAGATCTGG-3'
Avr-rev 5'-CACGCCTAGGCCGGTCGGTCTCGGGCCAC-3'
Nhe-rev 5'-GCGGCTAGCTGCTCGCCCATCGCGGGATGC-3'

The PCR included, in a 50 µl reaction, 5 µl of 10x *Pfu* polymerase buffer (Stratagene), 5 µl 10x z-dNTP mixture (2 mM dATP, 2 mM dCTP, 2 mM dTTP, 1 mM dGTP, 1 mM 7-deaza-GTP), 5 µl DMSO, 2 µl of each primer (10 µM), 1 µl of template DNA (0.1 µg/µl), and 1 µl of cloned *Pfu* polymerase (Stratagene). The PCR conditions were 95°C for 2 min., 25 cycles at 95°C for 30 sec., 60°C for 30 sec., and 72°C for 4

- 88 -

min., followed by 4 min. at 72°C and a hold at 0°C. The amplified DNA products and the Litmus vectors were cut with the appropriate restriction enzymes (*Bgl*III and *Avr*II or *Spe*I and *Nhe*I), and cloned into either pLitmus 28 or pLitmus38 (New England Biolabs), respectively, to generate the constructs designated pKOS60-37-4 and pKOS60-37-2, respectively.

Plasmid pKOS60-29-55 was again used as a template for PCR to amplify sequence 3' to the AT domain using the primers BsrXho-fwd and NsiAfl-rev:

BsrXho-fwd 5'-GATGTACAGCTCGAGTCGGCACGCCCCGGCCGCATC-3'

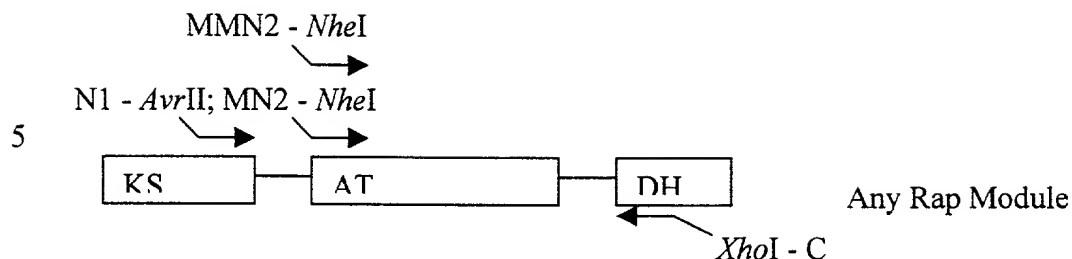
NsiAfl-rev 5'-CGACTCACTTAAGCCATGCATCC-3'

PCR conditions were as described above. The PCR fragment was cut with *Bsr*GI and *Afl*III, gel isolated, and ligated into pKOS60-37-4 cut with *Asp*718 and *Afl*III and inserted into pKOS60-37-2 cut with *Bsr*GI and *Afl*III, to give the plasmids pKOS60-39-1 and pKOS60-39-13, respectively. These two plasmids can be digested with *Avr*II and *Xho*I or *Nhe*I and *Xho*I, respectively, to insert heterologous AT domains specific for malonyl, methylmalonyl, ethylmalonyl, or other extender units.

Malonyl and methylmalonyl-specific AT domains were cloned from the rapamycin cluster using PCR amplification with a pair of primers that introduce an *Avr*II or *Nhe*I site at the 5' end and an *Xho*I site at the 3' end. The PCR conditions were as given above and the primer sequences were as follows:

RATN1 5'-ATCCTAGGCGGGCRGGYGTGTCGTCCTTCGG-3'
(3' end of Rap KS sequence and universal for malonyl and methylmalonyl CoA),
RATMN2 5'-ATGCTAGCCGCGCGTTCCCCGTCTTCGCGCG-3'
(Rap AT shorter version 5'- sequence and specific for malonyl CoA),
RATMMN2 5'-ATGCTAGCGGATTCGTCGGTGGTGTTCGCCGA-3'
(Rap AT shorter version 5'- sequence and specific for methylmalonyl CoA), and
RATC 5'-ATCTCGAGCCAGTASCGCTGGTGYTGGAAGG-3'
(Rap DH 5'- sequence and universal for malonyl and methylmalonyl CoA).

- 89 -



10 Because of the high sequence similarity in each module of the rapamycin cluster, each primer was expected to prime any of the AT domains. PCR products representing ATs specific for malonyl or methylmalonyl extenders were identified by sequencing individual cloned PCR products. Sequencing also confirmed that the chosen clones contained no cloning artifacts. Examples of hybrid modules with the rapamycin AT12 and AT13 domains are shown in a separate figure.

15 The *AvrII-XhoI* restriction fragment that encodes module 8 of the FK-520 PKS with the endogenous AT domain replaced by the AT domain of module 12 of the rapamycin PKS has the DNA sequence and encodes the amino acid sequence shown below. The AT of rap module 12 is specific for incorporation of malonyl units.

20 AGATCTGGCAGCTCGCCGAAGCGCTGCTGACGCTCGTCCGGGAGAGCACC 50
I W Q L A E A L L T L V R E S T
GCCGCCGTGCTCGGCCACGTGGGTGGCGAGGACATCCCCGCGACGGCGGC 100
A A V L G H V G G E D I P A T A A
GTTCAAGGACCTCGGCATCGACTCGCTCACCGCGGTCCAGCTGCGCAACG 150
25 F K D L G I D S L T A V Q L R N
CCCTCACCGAGGCGACCGGTGTGCGGCTGAACGCCACGGCGGTCTTCGAC 200
A L T E A T G V R L N A T A V F D
TTCCCGACCCGCGACGTGCTCGCCGGGAAGCTCGGCGACGAAGTACCGG 250
F P T P H V L A G K L G D E L T G
30 CACCCGCGCGCCCGTCTGTCGCCCGGACCGCGGCCACGGCCGGTGCACG 300
T R A P V V P R T A A T A G A H
ACGAGCCGCTGGCGATCGTGGGAATGGCCTGCCGGCTGCCCGGCGGGGTC 350
D E P L A I V G M A C R L P G G V
GCGTCACCCGAGGAGCTGTGGCACCTCGTGGCATCCGGCACCGACGCCAT 400
35 A S P E E L W H L V A S G T D A I
CACGGAGTTCCCGACGGACCGCGGCTGGGACGTCGACGCGATCTACGACC 450
T E F P T D R G W D V D A I Y D
CGGACCCCGACGCGATCGGCAAGACCTTCGTCCGGCACGGTGGCTTCCTC 500
P D P D A I G K T F V R H G G F L
40 ACCGGCGCGACAGGCTTCGACGCGGCGTTCTTCGGCATCAGCCGCGCGA 550
T G A T G F D A A F F G I S P R E
GGCCCTCGCATGGACCCGAGCAGCGGGTGCTCCTGGAGACGTCTGTGGG 600
A L A M D P Q Q R V L L E T S W
AGCGGTTCAAAGCGCCGGCATACCCCGGACTCGACCCGCGGCAGCGAC 650

- 90 -

E A F E S A G I T P D S T R G S D
ACCGGCGTGTTCGTCGGCGCCTTCTCCTACGGTTACGGCACCGGTGCGGA 700
T G V F V G A F S Y G Y G T G A D
CACCGACGGCTTCGGCGCGACCGGCTCGCAGACCAAGTGTGCTCTCCGGCC 750
5 T D G F G A T G S Q T S V L S G
GGCTGTGCTACTTCTACGGTCTGGAGGGTCCGGCGGTACGGTCGACACG 800
R L S Y F Y G L E G P A V T V D T
GCGTGTTCGTCGCTCGCTGGTGGCGCTGCACCAGGCCGGGCAGTCGCTGCG 850
A C S S S L V A L H Q A G Q S L R
10 CTCCGGCGAATGCTCGCTCGCCCTGGTCGGCGGGCGTCACGGTGATGGCGT 900
S G E C S L A L V G G V T V M A
CTCCCGGGCGGCTTCGTGGAGTTCTCCCGGCAGCGCGGCTCGCGCCGGAC 950
S P G G F V E F S R Q R G L A P D
GGCCGGGCGAAGGCGTTTCGGCGCGGGTGCAGGACGGCACGAGCTTCGCCGA 1000
15 G R A K A F G A G A D G T S F A E
GGGTGCCGGTGTGCTGCTCGTCGAGAGGCTCTCCGACGCCGAACGCAACG 1050
G A G V L I V E R L S D A E R N
GTCACACCGTCCTGGCGGTTCGTCCGTGGTTTCGGCGGTCAACCAGGATGGT 1100
G H T V L A V V R G S A V N Q D G
20 GCCTCCAACGGGCTGTGCGCGCCGAACGGGCGGTTCGACGAGCGGGTGAT 1150
A S N G L S A P N G P S Q E R V I
CCGGCAGGCCCTGGCCAACGCCGGGCTCACCCCGGGCGGACGTGGACGCCG 1200
R Q A L A N A G L T P A D V D A
TCGAGGCCCCACGGCACCGGCACAGGCTGGGCGACCCCATCGAGGCACAG 1250
25 V E A H G T G T R L G D P I E A Q
GCGGTACTGGCCACCTACGGACAGGAGCGCGCCACCCCTGCTGCTGGG 1300
A V L A T Y G Q E R A T P L L L G
CTCGCTGAAGTCCAACATCGGCCACGCCAGGCCGCGTCCGGCGTCGCCG 1350
S L K S N I G H A Q A A S G V A
30 GCATCATCAAGATGGTGCAGGCCCTCCGGCACGGGGAGCTGCCGCCGACG 1400
G I I K M V Q A L R H G E L P P T
CTGCACGCCGACGAGCCGTCGCCGACGTCGACTGGACGGCCGGCGCCGT 1450
L H A C D E P S P H V D W T A G A V
CGAACTGCTGACGTCGGCCCCGGCGTGGCCCCGAGACCGACCGGCCTAGGC 1500
35 E L L T S A R P W P E T D R P R
GGGCAGGCGTGTGCTCCTTCGGGATCAGTGGCACCAACGCCCACGTCATC 1550
R A G V S S F G I S G T N A H V I
CTGGAAGCGCACCCCCACTCAGCCTGCGGACAACGCGGTGATCGAGCG 1600
L E S A P P T Q P A D N A V I E R
40 GGCACCGGAGTGGGTGCCGTTGGTGATTTTCGGCCAGGACCCAGTCGGCTT 1650
A P E W V P L V I S A R T Q S A
TGA CTGAGCACGAGGGCCGGTTGCGTGCGTATCTGGCGGCGTCGCCCCGG 1700
L T E H E G R L R A Y L A A S P G
GTGGATATGCGGGCTGTGGCATCGACGCTGGCGATGACACGGTCGGTGTT 1750
45 V D M R A V A S T L A M T R S V F
CGAGCACCGTGCCGTGCTGCTGGGAGATGACACCGTCACCGGCACCGCTG 1800
E H R A V L L G D D T V T G T A
TGTCTGACCCTCGGGCGGTGTTTCGTCTTCCCGGGACAGGGGTGCGAGCGT 1850
V S D P R A V F V F P G Q G S Q R
50 GCTGGCATGGGTGAGGAAC TGCCCGCGGTTCCCGCTCTTCGCGCGGAT 1900
A G M G E E L A A A F P V F A R I
CCATCAGCAGGTGTGGGACCTGCTCGATGTGCCCGATCTGGAGGTGAACG 1950
H Q Q W D L L D V P D L E V N
AGACCGGTTACGCCCAGCCGCCCTGTTTCGCAATGCAGGTGGCTCTGTTC 2000

- 91 -

094016 03230 9104660

E T G Y A Q P A L F A M Q V A L F
GGGCTGCTGGAATCGTGGGGTGTACGACCGGACGCGGTGATCGGCCATTC 2050
G L L E S W G V R P D A V I G H S
5 GGTGGGTGAGCTTGGCGCTGCGTATGTGTCCGGGGTGTGGTCGTTGGAGG 2100
V G E L A A A Y V S G V W S L E
ATGCCTGCACTTTGGTGTGCGCGCGGGCTCGTCTGATGCAGGCTCTGCCC 2150
D A C T L V S A R A R L M Q A L P
GCGGGTGGGGTGTATGGTGTGCTCCCGGTCTCGGAGGATGAGGCCCGGGC 2200
A G G V M V A V P V S E D E A R A
10 CGTGCTGGGTGAGGGTGTGGAGATCGCCGCGGTCAACGGCCCGTCGTCGG 2250
V L G E G V E I A A V N G P S S
TGGTTCTCTCCGGTGTAGAGCCGCCGTGCTGCAGGCCGCGGAGGGGGTG 2300
V V L S G D E A A V L Q A A E G L
GGGAAGTGGACGCGGCTGGCGACCGACGCGTTCATTCCGCCCGTAT 2350
15 G K W T R L A T S H A F H S A R M
GGAACCCATGCTGGAGAGTTCCGGGCGGTGCGCGAAGGCCTGACCTACC 2400
E P M L E E F R A V A E G L T Y
GGACGCCGCGAGGTCTCCATGGCCGTTGGTGTATCAGGTGACCACCGCTGAG 2450
R T P Q V S M A V G D Q V T T A E
20 TACTGGGTGCGGCAGGTCCGGGACACGGTCCGGTTCGCGGAGCAGGTGGC 2500
Y W V R Q V R D T V R F G E Q V A
CTCGTACGAGGACGCCGTGTTCTGTCGAGCTGGGTGCCGACCGGTCACTGG 2550
S Y E D A V F V E L G A D R S L
CCCGCCTGGTGCAGCGGTGTGCGATGCTGCACGGCGACCACGAAATCCAG 2600
25 A R L V D G V A M L H G D H E I Q
GCCGCGATCGGCGCCCTGGCCACCTGTATGTCAACGGCGTCACGGTCGA 2650
A A I G A L A H L Y V N G V T V D
CTGGCCCCGCGCTCCTGGGCGATGCTCCGGCAACACGGGTGCTGGACCTTC 2700
W P A L L G D A P A T R V L D L
30 CGACATACGCCTTCCAGCACCAGCGCTACTGGCTCGAGTCGGCACGCCCCG 2750
P T Y A F Q H Q R Y W L E S A R P
GCCGCATCCGACGCGGGCCACCCCGTGTGGGCTCCGGTATCGCCCTCGC 2800
A A S D A G H P V L G S G I A L A
CGGGTCCGCGGGGCGGGTTCACGGGTTCGGTGCCGACCGGTGCGGACC 2850
35 G S P G R V F T G S V P T G A D
GCGCGGTGTTCTGTCGCCGAGTGGCGCTGGCCCGCGGACGCGGTGAC 2900
R A V F V A E L A L A A A D A V D
TGCGCCACGGTCGAGCGGCTCGACATCGCCTCCGTGCCCGGCCGCGGG 2950
C A T V E R L D I A S V P G R P G
40 CCATGGCCGGACGACCGTACAGACCTGGGTGACGAGCCGGCGGACGACG 3000
H G R T T V Q T W V D E P A D D
GCCGGCGCCGGTTCACCGTGCACACCCGACCGGCGACGCCCCGTGGACG 3050
G R R R F T V H T R T G D A P W T
CTGCACGCCGAGGGGGTGTGCGCCCCATGGCACGGCCCTGCCCGATGC 3100
45 L H A E G V L R P H G T A L P D A
GGCCGACGCCGAGTGGCCCCACCGGGCGCGGTGCCCGGACGGGCTGC 3150
A D A E W P P P G A V P A D G L
CGGGTGTGTGGCGCCGGGGGACCAGGTCTTCGCCGAGGCCGAGGTGGAC 3200
P G V W R R G D Q V F A E A E V D
50 GGACCGGACGGTTTCGTGGTGCACCCCGACCTGCTCGACGCGGTCTTCTC 3250
G P D G F V V H P D L L D A V F S
CGCGGTCCGGGACGGAAGCCGCCAGCCGGCCGGATGGCGGACCTGACGG 3300
A V G D G S R Q P A G W R D L T
TGCACGCGTCCGACGCCACCGTACTGCGCGCTGCCTACCCGGCGCACC 3350

- 92 -

V H A S D A T V L R A C L T R R T
GACGGAGCCATGGGATTGCGCGCCTTCGACGGCGCCGGCCTGCCGGTACT 3400
D G A M G F A A F D G A G L P V L
CACC CGGAGGCGGTGACGCTGCGGGAGGTGGCGTCACCGTCCGGCTCCG 3450
5 T A E A V T L R E V A S P S G S
AGGAGTCGGACGGCCTGCACCGGTTGGAGTGGCTCGCGGTGCCGAGGCG 3500
E E S D G L H R L E W L A V A E A
GTCTACGACGGTGACCTGCCCCGAGGGACATGTCTGATCACC GCCGCCCA 3550
V Y D G D L P E G H V L I T A A H
10 CCCCACGACCCCCGAGGACATAACCCACCCGCGCCACACCCGCGCCACCC 3600
P D D P E D I P T R A H T R A T
GCGTCCTGACCGCCCTGCAACACCTCACCACCACCGACCACACCTC 3650
R V L T A L Q H H L T T T D H T L
ATCGTCCACACCACCGACCCCGCGGGCGCCACCGTCACCGGCCTCAC 3700
15 I V H T T T D P A G A T V T G L T
CCGCAACGCCCCAGAACGAACACCCCCACCGCATCCGCCTCATCGAAACCG 3750
R T A Q N E H P H R I R L I E T
ACCACCCCCACACCCCCCTCCCCCTGGCCCAACTCGCCACCCTCGACCAC 3800
D H P H T P L P L A Q L A T L D H
20 CCCCACCTCCGCCTCACCACACACCTCCACCACCCCCACCTCACCCC 3850
P H L R L T H H T L H H P H L T P
CCTCCACACCACACCCACCCACCCACCCCTCAACCCCGAACACG 3900
L H T T T P P T T T P L N P E H
CCATCATCATCACC GGCGGTCCGGCACCCCTCGCCGGCATCCTCGCCCGC 3950
25 A I I I T G G S G T L A G I L A R
CACCTGAACCACCCCCACACCTACCTCCTCTCCCGACCCCCACCCCCGA 4000
H L N H P H T Y L L S R T P P P D
CGCCACCCCCGGCACCCACCTCCCCTGCGACGTGCGGACCCCCACCAAC 4050
A T P G T H L P C D V G D P H Q
30 TCGCCACCACCTCACCACATCCCCCAACCCCTCACC GCCCATCTTCCAC 4100
L A T T L T H I P Q P L T A I F H
ACCGCCGCCACCCCTCGACGACGGCATCCTCCACGCCCTCACCCCGACCG 4150
T A A T L D D G I L H A L T P D R
CCTCACCACCGTCCTCCACCCCAAGCCAACGCCGCTGGCACCTGCACC 4200
35 L T T V L H P K A N A A W H L H
ACCTCACCCAAAACCAACCCCTCACCACCTTCGTCTCTACTCCAGCGCC 4250
H L T Q N Q P L T H F V L Y S S A
GCCGCCGTCTCTGGCAGCCCCGGACAAGGAACTACGCCGCCGCCAACGC 4300
A A V L G S P G Q G N Y A A A N A
40 CTTCTCGACGCCCTCGCCACCCACCGCCACACCTCGGCCAACCCGCCA 4350
F L D A L A T H R H T L G Q P A
CCTCCATCGCCTGGGGCATGTGGCACACCACCAGCACCTCACC GGACAA 4400
T S I A W G M W H T T S T L T G Q
CTCGACGACGCCGACCGGGACCGCATCCGCCGCGGCGGTTTCCTCCCGAT 4450
45 L D D A D R D R I R R G G F L P I
CACGGACGACGAGGGCATGGGGATGCAT
T D D E G

The *AvrII-XhoI* restriction fragment that encodes module 8 of the FK-520 PKS
50 with the endogenous AT domain replaced by the AT domain of module 13 (specific for

- 93 -

methylmalonyl CoA) of the rapamycin PKS has the DNA sequence and encodes the amino acid sequence shown below.

AGATCTGGCAGCTCGCCGAAGCGCTGCTGACGCTCGTCCGGGAGAGCACC 50
Q L A E A L L T L V R E S T
5 GCCGCGGTGCTCGGCCACGTGGGTGGCGAGGACATCCCCGCGACGGCGGC 100
A A V L G H V G G E D I P A T A A
GTTCAAGGACCTCGGCATCGACTCGCTCACCGCGGTCCAGCTGCGCAACG 150
F K D L G I D S L T A V Q L R N
CCCTCACCGAGGCGACCGGTGTGCGGCTGAACGCCACGGCGGTCTTCGAC 200
10 A L T E A T G V R L N A T A V F D
TTCCCGACCCCGCACGTGCTCGCCGGGAAGCTCGGCGACGAACTGACCGG 250
F P T P H V L A G K L G D E L T G
CACCCGCGCGCCCGTCTGTCGCCCGGACCGCGGCCACGGCCGGTGCGCACG 300
T R A P V V P R T A A T A G A H
15 ACGAGCCGCTGGCGATCGTGGGAATGGCCTGCCGCTGCCCGGCGGGGTC 350
D E P L A I V G M A C R L P G G V
GCGTCACCCGAGGAGCTGTGGCACCTCGTGGCATCCGGCACCGACGCCAT 400
A S P E E L W H L V A S G T D A I
CACGGAGTTCGCGACGGACCGCGGCTGGGACGTGACGCGATCTACGACC 450
20 T E F P T D R G W D V D A I Y D
CGGACCCCGACGCGATCGGCAAGACCTTCGTCCGGCACGGTGGCTTCCTC 500
P D P D A I G K T F V R H G G F L
ACCGGCGCGACAGGCTTCGACGCGGCTTCTTCGGCATCAGCCCGCGCGA 550
T G A T G F D A A F F G I S P R E
25 GGCCCTCGCGATGGACCCGACGAGCGGGTGCTCCTGGAGACGTCTGTTGG 600
A L A M D P Q Q R V L L E T S W
AGGCGTTTCAAAGCGCCGGCATCACCCCGGACTCGACCCGCGGCAGCGAC 650
E A F E S A G I T P D S T R G S D
ACCGGCGTGTTCGTGCGGCCTTCTCCTACGGTTACGGCACCGGTGCGGA 700
30 T G V F V G A F S Y G Y G T G A D
CACCGACGGCTTCGGCGCGACCGGCTCGCAGACCACTGTGCTCTCCGGCC 750
T D G F G A T G S Q T S V L S G
GGCTGTCTGTACTTCTACGGTCTGGAGGGTCCGGCGGTACGGTTCGACACG 800
R L S Y F Y G L E G P A V T V D T
35 GCGTGTTCGTGCTGCTGGTGGCGCTGCACCAAGCCGGGCAGTCGCTGCG 850
A C S S S L V A L H Q A G Q S L R
CTCCGGCGAATGCTCGCTCGCCCTGGTGGCGGGCTCACGGTGATGGCGT 900
S G E C S L A L V G G V T V M A
CTCCCGGCGGCTTCGTGGAGTTCTCCCGGCAGCGCGGCTTCGCGCCGGAC 950
40 S P G G F V E F S R Q R G L A P D
GGCCGGGCGAAGCGTTCGGCGCGGGTGGCGACGGCACGAGCTTCGCCGA 1000
G R A K A F G A G A D G T S F A E
GGGTGCCGGTGTGCTGATCGTCGAGAGGCTCTCCGACGCCGAACGCAACG 1050
G A G V L I V E R L S D A E R N
45 GTCACACCGTCTGGCGGTGCTCCGTGGTTCGGCGGTCAACCAGGATGGT 1100
G H T V L A V V R G S A V N Q D G
GCCTCCAACGGGCTGTGCGGCCCGAACGGGCGGTGCGAGGAGCGGGTGAT 1150
A S N G L S A P N G P S Q E R V I
CCGGCAGGCCCTGGCCAACGCCGGGCTCACCCCGGCGGACGTGGACGCCG 1200
50 R Q A L A N A G L T P A D V D A
TCGAGGCCCCACGGCACCGGCACCAAGGCTGGGCGACCCCATCGAGGCACAG 1250
V E A H G T G T R L G D P I E A Q

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GCGGTACTGGCCACCTACGGACAGGAGCGCGCCACCCCTGCTGCTGGG 1300
A V L A T Y G Q E R A T P L L L G
CTCGCTGAAGTCCAACATCGGCCACGCCAGGCCGCTCCGGCGTCGCCG 1350
S L K S N I G H A Q A A S G V A
GCATCATCAAGATGGTGCAGGCCCTCCGGCACGGGGAGCTGCCGCCGACG 1400
G I I K M V Q A L R H G E L P P T
CTGCACGCCGACGAGCCGTCGCCGCACGTCGACTGGACGGCCGGCGCCGT 1450
L H A D E P S P H V D W T A G A V
CGAACTGCTGACGTCGGCCCCGGCCGTGGCCCCGAGACCGACCGGCCTAGGC 1500
E L L T S A R P W P E T D R P R
GGGCGGGCGTGTCTCTTCGGAGTCAGCGGCACCAACGCCACGTCATC 1550
R A G V S S F G V S G T N A H V I
CTGGAGAGCGCACCCCCGCTCAGCCCCGGAGGAGGCGCAGCCTGTTGA 1600
L E S A P P A Q P A E E A Q P V E
GACGCCGGTGGTGGCCTCGGATGTGCTGCCGCTGGTGATATCGGCAAGA 1650
T P V V A S D V L P L V I S A K
CCCAGCCCCCCTGACCGAACACGAAGACCGGCTGCGCGCTACCTGGCG 1700
T Q P A L T E H E D R L R A Y L A
GCGTCGCCCCGGGGCGGATATACGGGCTGTGGCATCGACGCTGGCGGTGAC 1750
A S P G A D I R A V A S T L A V T
ACGGTCGGTGTTCGAGCACCGCGCCGTACTCCTTGAGATGACACCGTCA 1800
R S V F E H R A V L L G D D T V
CCGGCACCGCGGTGACCGACCCCAGGATCGTGTTCCTTTCCCGGGCAG 1850
T G T A V T D P R I V F V F P G Q
GGGTGGCAGTGGCTGGGGATGGGCAGTGCAGTGCAGGATTCGTCGGTGGT 1900
G W Q W L G M G S A L R D S S V V
GTTCCGCCGAGCGGATGGCCGAGTGTGCGGCGGCGTTGCGCGAGTTCGTGG 1950
F A E R M A E C A A A L R E F V
ACTGGGATCTGTTCACGGTTCTGGATGATCCGGCGGTGGTGGACCGGGTT 2000
D W D L F T V L D D P A V V D R V
GATGTGGTCCAGCCCGCTTCCTGGGCGATGATGGTTTCCTGGCCGCGGT 2050
D V V Q G P A S G W A M M V S L A A V
GTGGCAGGCGCCGCTGTGCGCCGCGATGCGGTGATCGGCCATTTCGAGG 2100
W Q A A G V R P D A V I G H S Q
GTGAGATCGCCGAGCTTGTGTGGCGGTGCGGTGTCACTACGCGATGCC 2150
G E I A A A C V A G A V S L R D A
GCCCCGATCGTGACCTTGCGCAGCCAGGCGATCGCCCGGGGCTGGCGGG 2200
A R I V T L R S Q A I A R G L A G
CCGGGGCGCGATGGCATCCGTCGCCCTGCCGCGCAGGATGTCGAGCTGG 2250
R G A M A S V A L P A Q D V E L
TCGACGGGGCCTGGATCGCCGCCCCACAACGGGCCCCGCTCCACCGTGATC 2300
V D G A W I A A H N G P A S T V I
GCGGGCACCCCGGAAGCGGTGACCATGTCTCACCCTCATGAGGCACA 2350
A G T P E A V D H V L T A H E A Q
AGGGGTGCGGGTGCAGCGGATCACCGTCGACTATGCCTCGCACACCCCGC 2400
G V R V R R I T V D Y A S H T P
ACGTCGAGCTGATCCGCGACGAACACTCGACATCACTAGCGACAGCAGC 2450
H V E L I R D E L L D I T S D S S
TCGACAGACCCCGCTCGTGCCGTGGCTGTGACCGTGGACGGCACCTGGGT 2500
S Q T P L V P W L S T V D G T W V
CGACAGCCCGCTGGACGGGGAGTACTGGTACCGGAACCTGCGTGAACCGG 2550
D S P L D G E Y W Y R N L R E P
TCGGTTTCCACCCCGCGTCAGCCAGTTGCAGGCCAGGGCGACACCGTG 2600
V G F H P A V S Q L Q A Q G D T V

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TTCGTCGAGGTCAGCGCCAGCCCGGTGTTGTTGCAGGCGATGGACGACGA 2650
F V E V S A S P V L L Q A M D D D
TGTCGTCACGGTTGCCACGCTGCGTCGTGACGACGGCGACGCCACCCGGA 2700
V V T V A T L R R D D G D A T R
TGCTCACCGCCCTGGCACAGGCCTATGTCCACGGCGTCACCGTCGACTGG 2750
M L T A L A Q A Y V H G V T V D W
CCCGCCATCCTCGGCACCACCACAACCCGGGTACTGGACCTTCCGACCTA 2800
P A I L G T T T T T R V L D L P T Y
CGCCTTCCAACACCAGCGGTACTGGCTCGAGTCGGCACGCCCCGGCCGCAT 2850
A F Q H Q R Y W L E S A R P A A
CCGACGCGGGGCCACCCCGTGCTGGGCTCCGGTATCGCCCTCGCCGGGTGCG 2900
S D A G H P V L G S G I A L A G S
CCGGGCGGGGTGTTACGGGTTCCGTGCCGACCGGTGCGGACCGCGCGGT 2950
P G R V F T G S V P T G A D R A V
GTTTCGTGCGCGAGCTGGCGCTGGCGCGCGGACGCGGTGCGACTGCGCCA 3000
F V A E L A L A A A D A V D C A
CGGTGCGAGCGGCTCGACATCGCCTCCGTGCCCCGGCCGGCCGGCCATGGC 3050
T V E R L D I A S V P G R P G H G
CGGACGACCGTACAGACCTGGGTGCGACGAGCCGGCGGACGACGGCCGGCG 3100
R T T V Q T W V D E P A D D G R R
CCGGTTCACCGTGACACCCGACCGGGCGACGCCCCGTGGACGCTGCACG 3150
R F T V H T R T G D A P W T L H
CCGAGGGGGTGCTGCGCCCCCATGGCACGGCCCTGCCCGATGCGGGCCGAC 3200
A E G V L R P H G T A L P D A A D
GCCGAGTGGCCCCACCGGGCGCGGTGCCCGCGGACGGGCTGCCGGGTGT 3250
A E W P P P G A V P A D G L P G V
GTGGCGCGGGGGGACCAGGTCTTCGCCGAGGCCGAGGTGGACGGACCGG 3300
W R R G G D Q V F A E A E V D G P
ACGGTTTTCGTGGTGACCCCGACCTGCTCGACGCGGTCTTCTCCGCGGTC 3350
D G F V V H P D L L D A V F S A V
GGCGACGGAAGCCGCCAGCCGGCCGGATGGCGCGACCTGACGGTGACGC 3400
G D G S R Q P A G W R D L T V H A
GTCGGACGCCACCGTACTGCGCGCCTGCCTACCCGGCGCACCGACGGAG 3450
S D A T V L R A C L T R R T D G
CCATGGGATTGCGCGCTTCGACGGCGCCGGCCTGCCGGTACTCACCGCG 3500
A M G F A A F D G A G L P V L T A
GAGGCGGTGACGCTGCGGGAGGTGGCGTCACCGTCCGGCTCCGAGGAGTC 3550
E A V T L R E V A S P S G S E E S
GGACGGCCTGCACCGGTTGGAGTGGCTCGCGGTGCGCGAGGCGGTCTACG 3600
D G L H R L E W L A V A E A V Y
ACGGTGACCTGCCCGAGGGACATGTCCTGATCACCGCCGCCACCCCGAC 3650
D G D L P E G H V L I T A A H P D
GACCCCGAGGACATACCCACCCGCGCCACACCCGCGCCACCCGCGTCCT 3700
D P E D I P T R A H T R A T R V L
GACCGCCCTGCAACACCACCTCACCACCACCGACCACACCCTCATCGTCC 3750
T A L Q H L T T T D H T L I V
ACACCACCACCGACCCCGCGGGCGCCACCGTCACCGGCCTACCCGCACC 3800
H T T T D P A G A T V T G L T R T
GCCCAGAACGAACACCCCCACCGCATCCGCTCATCGAAACCGACCACCC 3850
A Q N E H P H R I R L I E T D H P
CCACACCCCCCTCCCCCTGGCCCAACTCGCCACCCTCGACCACCCCCACC 3900
H T P L P L A Q L A T L D H P H
TCGCGCTCACCCACCACACCCTCCACCACCCCCACCTACCCCCCTCCAC 3950
L R L T H H T L H H P H L T P L H

- 96 -

ACCACCACCCACCCACCCACCCACCCCTCAACCCCGAACACGCCATCAT 4000
T T T P P T T T P L N P E H A I I
CATCACCGGGCGGCTCCGGCACCTCGCGGCATCTCGCCCGCCACCTGA 4050
I T G G S G T L A G I L A R H L
5 ACCACCCCCACACCTACCTCCTCTCCCGCACCCACCCCGACGCCACC 4100
N H P H T Y L L S R T P P P D A T
CCCGGCACCCACCTCCCCTGCGACGTGCGGCGACCCCACTCGCCAC 4150
P G T H L P C D V G D P H Q L A T
CACCCTCACCCACATCCCCCAACCCCTCACCGCCATCTTCCACACCGCCG 4200
10 T L T H I P Q P L T A I F H T A
CCACCCTCGACGACGGCATCTCCACGCCCTCACCCCGACCGCCTCACC 4250
A T L D D G I L H A L T P D R L T
ACCGTCTCCACCCCAAAGCCAACGCCGCTGGCACCTGCACCACCTCAC 4300
T V L H P K A N A A W H L H H L T
15 CCAAAACCAACCCCTCACCCACTTCGTCCTCTACTCCAGCGCCGCGCCG 4350
Q N Q P L T H F V L Y S S A A A
TCCTCGGCAGCCCCGACAAGGAACTACGCCGCGCCAACGCCTTCCTC 4400
V L G S P G Q G N Y A A A N A F L
GACGCCCTCGCCACCCACCGCCACACCCTCGGCCAACCCGCCACCTCCAT 4450
20 D A L A T H R H T L G Q P A T S I
CGCCTGGGGCATGTGGCACACCACGACCCCTCACCGGACAACCTCGACG 4500
A W G M W H T T S T L T G Q L D
ACGCCGACCGGGACCGCATCCGCCGCGGCGGTTTCCTCCCGATCACGGAC 4550
D A D R D R I R R G G F L P I T D
25 GACGAGGGCATGGGGATGCAT
D E G

The *NheII-XhoI* restriction fragment that encodes module 8 of the FK-520 PKS
with the endogenous AT domain replaced by the AT domain of module 12 (specific for
malonyl CoA) of the rapamycin PKS has the DNA sequence and encodes the amino acid
sequence shown below.

AGATCTGGCAGCTCGCCGAAGCGCTGCTGACGCTCGTCCGGGAGAGCACC 50
Q L A E A L L T L V R E S T
GCCGCCGTGCTCGGCCACGTGGGTGGCGAGGACATCCCCGCGACGGCGGC 100
35 A A V L G H V G G E D I P A T A A
GTTCAAGGACCTCGGCATCGACTCGCTCACCGCGGTCCAGCTGCGCAACG 150
F K D L G I D S L T A V Q L R N
CCCTCACCGAGGCGACCGGTGTGCGGCTGAACGCCACGGCGGTCTTCGAC 200
A L T E A T G V R L N A T A V F D
40 TTCCCGACCCCGCACGTGCTCGCCGGAAGCTCGGCGACGAACTGACCGG 250
F P T P H V L A G K L G D E L T G
CACCCGCGCGCCCGTCTGTGCCCCGACCGCGGCCACGGCCGGTGCACG 300
T R A P V V P R T A A T A G A H
ACGAGCCGCTGGCGATCGTGGGAATGGCCTGCCGGCTGCCCGGCGGGGTC 350
45 D E P L A I V G M A C R L P G G V
GCGTCACCCGAGGAGCTGTGGCACCTCGTGGCATCCGGCACCGACGCCAT 400
A S P E E L W H L V A S G T D A I
CACGGAGTTCCCGACGGACCGCGGCTGGGACGTCGACCGATCTACGACC 450
T E F P T D R G W D V D A I Y D
50 CGGACCCCGACGCGATCGGCAAGACCTTCGTCCGGCACGGTGGCTTCCTC 500

- 97 -

P D P D A I G K T F V R H G G F L
ACCGGCGCGACAGGCTTCGACGCGGCGTTCTTCGGCATCAGCCCGCGCGA 550
T G A T G F D A A F F G I S P R E
GGCCCTCGCGATGGACCCGCGAGCAGCGGGTGCTCCTGGAGACGTCGTGGG 600
5 A L A M D P Q Q R V L L E T S W
AGGCGTTCGAAAGCGCCGGCATCACCCCGGACTCGACCCGCGGCAGCGAC 650
E A F E S A G I T P D S T R G S D
ACCGGCGTGTTCGTCGGCGCCTTCTCCTACGGTTACGGCACCGGTGCGGA 700
T G V F V G A F S Y G Y G T G A D
10 CACCGACGGCTTCGGCGCGACCGGCTCGCAGACCAAGTGTGCTCTCCGGCC 750
T D G F G A T G S Q T S V L S G
GGGTGTCGTACTTCTACGGTCTGGAGGGTCCGGCGGTACGGTTCGACACG 800
R L S Y F Y G L E G P A V T V D T
GCGTGTTCGTCGTCGCTGGTGGCGCTGCACCAGGCCGGGCAGTCGCTGCG 850
15 A C S S S L V A L H Q A G Q S L R
CTCCGGCAATGCTCGCTCGCCCTGGTTCGGCGGCGTCACGGTGATGGCGT 900
S G E C S L A L V G G V T V M A
CTCCCGGCGGCTTCGTGGAGTTCTCCCGGCAGCGCGCCTCGCGCCGGAC 950
S P G G F V E F S R Q R G L A P D
20 GGCCGGGCGAAGGCGTTCGGCGCGGGTTCGGACGGCACGAGCTTCGCCGA 1000
G R A K A F G A G A D G T S F A E
GGGTGCCGGTGTGCTGATCGTCGAGAGGCTCTCCGACGCCGAACGCAACG 1050
G A G V L I V E R L S D A E R N
GTCACACCGTCTGGCGGTCTGCTGGTTCGGCGGTCAACCAGGATGGT 1100
25 G H T V L A V V R G S A V N Q D G
GCCTCCAACGGGCTGTGCGGCGCCGAACGGGCGGTCGCGAGGAGCGGGTGAT 1150
A S N G L S A P N G P S Q E R V I
CCGGCAGGCCCTGGCCAACGCCGGGCTCACCCCGGCGGACGTGGACGCCG 1200
R Q A L A N A G L T P A D V D A
30 TCGAGGCCCCACGGCACCGGCACAGGCTGGGCGACCCCATCGAGGCACAG 1250
V E A H G T G T R L G D P I E A Q
GCGGTACTGGCCACCTACGGACAGGAGCGCGCCACCCCTGCTGCTGGG 1300
A V L A T Y G Q E R A T P L L L G
CTCGCTGAAGTCCAACATCGGCCACGCCAGGCCGCGTCCGGCGTCGCCG 1350
35 S L K S N I G H A Q A S G V A
GCATCATKAAGATGGTGACGGCCCTCCGGCACGGGAGCTGCCGCCGACG 1400
G I I K M V Q A L R H G E L P P T
CTGCACGCCGACGAGCCGTCGCCGCACGTCGACTGGACGGCCGGCGCCGT 1450
L H A D E P S P H V D W T A G A V
40 CGAACTGCTGACGTCGGCCCCGGCGTGGCCCCGAGACCGACCGGCCACGGC 1500
E L L T S A R P W P E T D R P R
GTGCCCGCTCTCCTCGTTCGGGGTGAGCGGCACCAACGCCACGTCATC 1550
R A A V S S F G V S G T N A H V I
CTGGAGCCCGACCGGTAACGAGACGCCCGCGGCATCGCCTTCCGGTGA 1600
45 L E A G P V T E T P A A S P S G D
CCTTCCCCTGCTGGTGTGCGGCACGCTCACCGGAAGCGCTCGACGAGCAGA 1650
L P L L V S A R S P E A L D E Q
TCCGCCGACTGCGCGCTACCTGGACACCACCCCGGACGTCGACCGGGTG 1700
I R R L R A Y L D T T P D V D R V
50 GCCGTGGCACAGACGCTGGCCCCGGCGCACACTTCGCCACCGCGCCGT 1750
A V A Q T L A R R T H F A H R A V
GCTGCTCGGTGACACCGTCATCACCACACCCCGCGGACCGGCCGACG 1800
L L G D T V I T T P P A D R P D
AACTCGTCTTCGTCTACTCCGGCCAGGGCACCCAGCATCCCGCGATGGGC 1850

[illegible][illegible]

- 99 -

R R G D Q V F A E A E V D G P D G
TTCGTGGTGCACCCCGACCTGCTCGACGCGGTCTTCTCCGCGGTCTGGCGA 3250
F V V H P D L L D A V F S A V G D
CGGAAGCCGCCAGCCGCGCGGATGGCGCGACCTGACGGTGCACGCGTCGG 3300
5 G S R Q P A G W R D L T V H A S
ACGCCACCGTACTGCGCGCCTGCCTCACCCGGCGCACCGACGGAGCCATG 3350
D A T V L R A C L T R R T D G A M
GGATTGCGCGCCTTCGACGGCGCGGCTGCCGGTACTCACCGCGGAGGC 3400
G F A A F D G A G L P V L T A E A
10 GGTGACGCTGCGGGAGGTGGCGTCACCGTCCGGCTCCGAGGAGTCGGACG 3450
V T L R E V A S P S G S E E S D
GCCTGCACCGGTTGGAGTGGCTCGCGGTGCGCGAGGCGGTCTACGACGGT 3500
G L H R L E W L A V A E A V Y D G
GACCTGCCCCGAGGGACATGTCTGATCACCGCGCGCCACCCCGACGACCC 3550
15 D L P E G H V L I T A A H P D D P
CGAGGACATACCCACCCGCGCCACACCCGCGCCACCCGCGTCTGACCG 3600
E D I P T R A H T R A T R V L T
CCCTGCAACACCACTCACCAACACCGACCAACACCTCATCGTCCACACC 3650
A L Q H H L T T T D H T L I V H T
20 ACCACCGACCCCGCGCGGCCACCGTCACCGGCCTCACCCGCACCGCCCA 3700
T T D P A G A T V T G L T R T A Q
GAACGAACACCCCCACCGCATCCGCTCATCGAAACCGACACCCCCACA 3750
N E H P H R I R L I E T D H P H
CCCCCTCCCCCTGGCCCCAACTCGCCACCTCGACCACCCCACTCCGC 3800
25 T P L P L A Q L A T L D H P H L R
CTCACCAACCAACCTCCACCAACCCCACTCACCCCTCCACACCAC 3850
L T H H T L H H P H L T P L H T T
CACCCCAACCAACCAACCCCTCAACCCGAACACGCCATCATCATCA 3900
T P P T T T P L N P E H A I I I
30 CCGGCGGCTCCGGCACCTCGCGGCATCTCGCCCGCCACCTGAACCAC 3950
T G G S G T L A G I L A R H L N H
CCCCACACCTACCTCTCTCCCGCACCCCAACCCCGACGCCACCCCGG 4000
P H T Y L L S R T P P P D A T P G
CACCCACCTCCCCTGCGACGTGCGCGACCCCACTCGCCACCAACC 4050
35 T H L P C D V G D P H Q L A T T
TCACCCACATCCCCCAACCCCTCACCGCATCTTCCACACCGCGCCACC 4100
L T H I P Q P L T A I F H T A A T
CTCGACGACGGCATCTCCACGCCCTCACCCCGACCGCTCACCACCGT 4150
L D D G I L H A L T P D R L T T V
40 CCTCCACCCCAAGCCAACGCGCCTGGCACCTGCACCACCTCACCCAAA 4200
L H P K A N A A W H L H H L T Q
ACCAACCCCTCACCACTTCGTCTCTACTCCAGCGCGCGCGTCTCTC 4250
N Q P L T H F V L Y S S A A A V L
GGCAGCCCCGACAAGGAACTACGCCGCCCAACGCCTTCTCGACGC 4300
45 G S P G Q G N Y A A A N A F L D A
CCTCGCCACCCACCGCCACACCTCGGCCAACCCGCCACCTCCATCGCCT 4350
L A T H R H T L G Q P A T S I A
GGGGCATGTGGCACACCACAGCACCTCACCGGACAACCTCGACGACGCC 4400
W G M W H T T S T L T G Q L D D A
50 GACCGGGACCGCATCCGCCGCGCGGTTTCTCCCGATCACGGACGACGA 4450
D R D R I R R G G F L P I T D D E
GGGCATGGGGATGCAT
G

FILED OCT 23 2006

- 100 -

The *NheII-XhoI* restriction fragment that encodes module 8 of the FK-520 PKS with the endogenous AT domain replaced by the AT domain of module 13 (specific for methylmalonyl CoA) of the rapamycin PKS has the DNA sequence and encodes the amino acid sequence shown below.

```

5  AGATCTGGCAGCTCGCCGAAGCGCTGCTGACGCTCGTCCGGGAGAGCACC 50
   Q L A E A L L T L V R E S T
   GCCGCCGTGCTCGGCCACGTGGGTGGCGAGGACATCCCCGCGACGGCGGC 100
   A A V L G H V G G E D I P A T A A
   GTTCAAGGACCTCGGCATCGACTCGCTCACC GCGGTCCAGCTGCGCAACG 150
10  F K D L G I D S L T A V Q L R N
   CCCTCACCGAGGCGACCGGTGTGCGGCTGAACGCCACGGCGGTCTTCGAC 200
   A L T E A T G V R L N A T A V F D
   TTCCCGACCCCGCACGTGCTCGCCGGGAAGCTCGGCGACGAAGTACCGG 250
   F P T P H V L A G K L G D E L T G
15  CACCCGCGCGCCCGTCTGTCGCCCCGACCGCGGCCACGGCCGGTGCGCACG 300
   T R A P V V P R T A A T A G A H
   ACGAGCCGCTGGCGATCGTGGGAATGGCCTGCCGGCTGCCCGCGGGGTC 350
   D E P L A I V G M A C R L P G G V
   GCGTCACCCGAGGAGCTGTGGCACCTCGTGGCATCCGGCACCGACGCCAT 400
20  A S P E E L W H L V A S G T D A I
   CACGGAGTTCCCGACGGACCGCGGCTGGGACGTGACGCGATCTACGACC 450
   T E F P T D R G W D V D A I Y D
   CGGACCCCGACGCGATCGGCAAGACCTTCGTCCGGCACGGTGGCTTCCTC 500
   P D P D A I G K T F V R H G G F L
25  ACCGGCGCGACAGGCTTCGACGCGGCGTTCTTCGGCATCAGCCCGCGCGA 550
   T G A T G F D A A F F G I S P R E
   GGCCCTCGCGATGGACCCGACGAGCGGGTGCTCCTGGAGACGTCTGGG 600
   A L A M D P Q Q R V L L E T S W
   AGGCGTTTCGAAAGCGCCGGCATCACCCCGGACTCGACCCGCGGCAGCGAC 650
30  E A F E S A G I T P D S T R G S D
   ACCGGCGTGTTCGTGCGGCCTTCTCCTACGGTTACGGCACCGGTGCGGA 700
   T G V F V G A F S Y G Y G T G A D
   CACCGACGGCTTCGGCGCGACCGGCTCGCAGACCAAGTGTGCTCTCCGGCC 750
   T D G F G A T G S Q T S V L S G
35  GGCTGTCTGTAATCTACGGTCTGGAGGGTCCGGCGGTACGGTCGACACG 800
   R L S Y F Y G L E G P A V T V D T
   GCGTGTTCGTGCTCGCTGGTGGCGCTGCACCAGGCCGGGACGTGCTGCG 850
   A C S S S L V A L H Q A G Q S L R
   CTCCGGCGAATGCTCGCTCGCCCTGGTCGGCGGCGTCACGGTGATGGCGT 900
40  S G E C S L A L V G G V T V M A
   CTCCCGGCGGCTTCGTGGAGTTCTCCCGGACGCGCGGCTCGCGCCGGAC 950
   S P G G G F V E F S R Q R G L A P D
   GGCCGGGCGAAGGCTTCGGCGCGGTCGCGACGGCACGAGCTTCGCCGA 1000
   G R A K A F G A G A D G T S F A E
45  GGGTGCCGGTGTGCTGATCGTCGAGAGGCTCTCCGACGCCGAACGCAACG 1050
   G A G V L I V E R L S D A E R N
   GTCACACCGTCTGGCGGTGCTCCGTGGTTCGGCGGTCAACCAGGATGGT 1100
   G H T V L A V V R G S A V N Q D G
   GCCTCCAACGGGCTGTGCGCGCCGAACGGGCGGTCGCGAGAGCGGGTGAT 1150
50  A S N G L S A P N G P S Q E R V I

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- 101 -

CCGGCAGGCCCTGGCCAACGCCGGGCTACCCCGGCGGACGTGGACGCCG 1200
R Q A L A N A G L T P A D V D A
TCGAGGCCACCGGCACCGGCACCGGCTGGGCGACCCCATCGAGGCACAG 1250
V E A H G T G T R L G D P I E A Q
5 GCGGTACTGGCCACCTACGGACAGGAGCGCGCCACCCCCCTGCTGCTGGG 1300
A V L A T Y G Q E R A T P L L L G
CTCGCTGAAGTCCAACATCGGCCACGCCAGGCCGCGTCCGGCGTCCGCCG 1350
S L K S N I G H A Q A A S G V A
GCATCATCAAGATGGTGCAGGCCCTCCGGCACGGGGAGCTGCCGCCGACG 1400
10 G I I K M V Q A L R H G E L P P T
CTGCACGCCGACGAGCCGTCGCCGCACGTCGACTGGACGGCCGGCGCCGT 1450
L H A D E P S P H V D W T A G A V
CGAAGTGTGACGTCGGCCCCGGCCGTGGCCCCGAGACCGACCGGCCACGGC 1500
E L L T S A R P W P E T D R P R
15 GTGCCGCCGTCTCCTCGTTCGGGGTGAGCGGCACCAACGCCACGTCATC 1550
R A A V S S F G V S G T N A H V I
CTGGAGGCCCGACCGGTAACGGAGACGCCCGCGGCATCGCCTTCCGGTGA 1600
L E A G P V T E T P A A S P S G D
20 CCTTCCCCTGCTGGTGTGCGGCACGCTACCGGAAGCGCTCGACGAGCAGA 1650
L P L L V S A R S P E A L D E Q
TCCGCGACTGCGCGCCTACCTGGACACACCCCGGACGTCGACCGGGTG 1700
I R R L R A Y L D T T P D V D R V
GCCGTGGCACAGACGCTGGCCCCGGCGCACACTTCGCCCCACCGCGCCGT 1750
A V A Q T L A R R T H F A H R A V
25 GCTGCTCGGTGACACCGTCATCACCACACCCCCCGCGGACCGGCCGACG 1800
L L G D T V I T T P P A D R P D
AACTCGTCTTCGTCTACTCCGGCCAGGGCACCCAGCATCCCGCGATGGGC 1850
E L V F V Y S G Q G T Q H P A M G
GAGCAGCTAGCCGATTCGTGCGGTGGTGTTCGCCGAGCGGATGGCCGAGTG 1900
30 E Q L A D S S V V F A E R M A E C
TGCGGCGCGGTTGCGCGAGTTCGTGGACTGGGATCTGTTACGGTTCTGG 1950
A A A L R E F V D W D L F T V L
ATGATCCGGCGGTGGTGGACCGGGTTGATGTGGTCCAGCCCGCTTCTGG 2000
D D P A V V D R V D V V Q P A S W
35 GCGATGATGGTTTCCCTGGCCGCGGTGTGGCAGGCGGCCGCTGTGCGGCC 2050
A M M V S L A A V W Q A A G V R P
GGATGCGGTGATCGGCCATTCGACGGGTGAGATCGCCGACGCTTGTGTGG 2100
D A V I G H S Q G E I A A A C V
CGGGTGCGGTGTCACTACGCGATGCCGCCCGGATCGTGACCTTGCGCAGC 2150
40 A G A V S L R D A A R I V T L R S
CAGGCGATCGCCCGGGGCTGGCGGGCGGGGCGCGATGGCATCCGTCGC 2200
Q A I A R G L A G R G A M A S V A
CCTGCCCGCGCAGGATGTCGAGCTGGTTCGACGGGGCTGGATCGCCGCC 2250
L P A Q D V E L V D G A W I A A
45 ACAACGGGCCCCCTCCACCGTGATCGCGGGCACCCCGGAAGCGGTTCGAC 2300
H N G P A S T V I A G T P E A V D
CATGTCCTCACCGCTCATGAGGCACAAGGGGTGCGGGTGCGGCGGATCAC 2350
H V L T A H E A Q G V R V R R I T
CGTCGACTATGCCTCGCACACCCCGCACGTCGAGCTGATCCGCGACGAAC 2400
50 V D Y A S H T P H V E L I R D E
TACTCGACATCACTAGCGACAGCAGCTCGCAGACCCCGCTCGTGCCGTGG 2450
L L D I T S D S S S Q T P L V P W
CTGTCGACCGTGGACGGCACCTGGGTCGACAGCCCGCTGGACGGGGAGTA 2500
L S T V D G T W V D S P L D G E Y

- 102 -

CTGGTACCGGAACCTGCGTGAACCGGTCGGTTTCCACCCCGCCGTCAGCC 2550
W Y R N L R E P V G F H P A V S
AGTTGCAGGCCCCAGGGCGACACCGTGTTCGTCGAGGTCAGCGCCAGCCCCG 2600
Q L Q A Q G D T V F V E V S A S P
5 GTGTTGTTGCAGGCGATGGACGACGATGTCGTCACGGTTGCCACGCTGCG 2650
V L L Q A M D D D V V T V A T L R
TCGTGACGACGGCGACGCCACCCGGATGCTCACC GCCCTGGCACAGGCCT 2700
R D D G D A T R M L T A L A Q A
ATGTCCACGGCGTCACCGTCGACTGGCCCCCATCCTCGGCACCACCACA 2750
10 Y V H G V T V D W P A I L G T T T
ACCCGGGTACTGGACCTTCCGACCTACGCCTTCCAACACCAGCGGTACTG 2800
T R V L D L P T Y A F Q H Q R Y W
GCTCGAGTCGGCAGCCCCGGCCGATCCGACGCGGGCCACCCCGTGTCTGG 2850
L E S A R P A A S D A G H P V L
15 GCTCCGGTATCGCCCTCGCCGGGTGCGCGGGCCGGGTGTTACGGGTTCC 2900
G S G I A L A G S P G R V F T G S
GTGCCGACCGGTGCGGACCGCGCGGTGTTTCGTCGCCGAGCTGGCGCTGGC 2950
V P T G A D R A V F V A E L A L A
CGCCGCGGACGCGGTGCGACTGCGCCACGGTCGAGCGGCTCGACATCGCCT 3000
20 A A D A V D C A T V E R L D I A
CCGTGCCCCGGCCGGCCGGCCATGGCCGGACGACCGTACAGACCTGGGTC 3050
S V P G R P P G H G R T T V Q T W V
GACGAGCCGGGACGACGGCCGGCCGGTTACCGTGCACACCCGCAC 3100
D E P A D D G R R R F T V H T R T
25 CGGCGACGCCCCGTGGACGCTGCACGCCGAGGGGTGCTGCGCCCCCATG 3150
G D A P W T L H A E G V L R P H
GCACGGCCCTGCCGATGCGGCCGACGCCGAGTGGCCCCACCGGGCGCG 3200
G T A L P D A A D A E W P P P G A
GTGCCCCGCGACGGGCTGCCGGGTGTGTGGCGCCGGGGGACCAGGTCTT 3250
30 V P A D G L P G V W R R G D Q V F
CGCCGAGGCCGAGGTGGACGGACCGGACGGTTTCGTGGTGCACCCCGACC 3300
A E A E V D G P D G F V V H P D
TGCTCGACGCGGTCTTCTCCGCGGTGCGCGACGGAAGCCGCCAGCCGGCC 3350
L L D A V F S A V G D G S R Q P A
35 GGATGGCGCGACCTGACGGTGCACGCGTCGGACGCCACCGTACTGCGCGC 3400
G W R D L T V H A S D A T V L R A
CTGCCTACCCGGCGCACCGACGGAGCCATGGGATTGCGCGCCTTCGACG 3450
C L T R R T D G A M G F A A F D
GCGCCGGCCTGCCGGTACTCACCGCGGAGGCGGTGACGCTGCGGGAGGTG 3500
40 G A G L P V L T A E A V T L R E V
GCGTCACCGTCCGGCTCCGAGGAGTCGGACGGCCTGCACCGTTGGAGTG 3550
A S P S G S E E S D G L H R L E W
GCTCGCGGTGCGCGAGGCGGTCTACGACGGTGACCTGCCCGAGGGACATG 3600
L A V A E A V Y D G D L P E G H
45 TCCTGATCACCGCCGCCCCACCCCGACGACCCCGAGGACATACCCACCCGC 3650
V L I T A A H P D D P E D I P T R
GCCACACCCGCGCCACCCGCGTCTGACCGCCCTGCAACACCACCTCAC 3700
A H T R A T R V L T A L Q H H L T
CACCACCGACCACACCTCATCGTCCACACCACCGACCCCGCCGGCG 3750
50 T T D H T L I V H T T T D P A G
CCACCGTACCGGCCTCACCCGCACCGCCAGAACGAACACCCCCACCGC 3800
A T V T G L T R T A Q N E H P H R
ATCCGCCTCATCGAAACCGACACCCCCACACCCCTCCCCCTGGCCCA 3850
I R L I E T D H P H T P L P L A Q

- 103 -

ACTCGCCACCCTCGACCACCCCCACCTCCGCCTCACCACCCACACCCCTCC 3900
L A T L D H P H L R L T H H T L
ACCACCCCCACCTCACCCTCCACACCACCCACCCACCCACCCACC 3950
H H P H L T P L H T T T P P T T T
5 CCCCTCAACCCGAACACGCCATCATCATACCGGCGGCTCCGGCACCCCT 4000
P L N P E H A I I I T G G S G T L
CGCCGGCATCTCGCCCGCCACCTGAACACCCCCACACCTACCTCCTCT 4050
A G I L A R H L N H P H T Y L L
CCCACACCCACCCCGGACGCCACCCCGGCACCCACCTCCCTGCGAC 4100
10 S R T P P P D A T P G T H L P C D
GTCGGCGACCCCACTCGCCACCACCCCTCACCACATCCCCCAACC 4150
V G D P H Q L A T T L T H I P Q P
CCTCACCGCCATCTTCCACACCGCGCCACCCCTCGACGACGGCATCCTCC 4200
L T A I F H T A A T L D D G I L
15 ACGCCCTCACCCTGACCGCCTCACCACCGTCTCCACCCAAAGCCAAC 4250
H A L T P D R L T T V L H P K A N
GCCGCCTGGCACCTGCACCACCTACCCAAAACCAACCCCTCACCCTT 4300
A A W H L H H L T Q N Q P L T H F
CGTCCTCTACTCCAGCGCGCGCGCTCCTCGGCAGCCCCGGACAAGGAA 4350
20 V L Y S S A A A V L G S P G Q G
ACTACGCCGCGCCCAACGCCTTCTCGACGCCCTCGCCACCCACCGCCAC 4400
N Y A A A N A F L D A L A T H R H
ACCTCGGCCAACCGCCACCTCCATCGCTGGGGCATGTGGCACACCAC 4450
T L G Q P A T S I A W G M W H T T
25 CAGACCTCACCAGCAACTCGACGACGCGGACCGGGACCGCATCCGCC 4500
S T L T G Q L D D A D R D R I R
GCGGCGGTTTCTCCGATCACGACGACGAGGGCATGGGGATGCAT
R G G F L P I T D D E G

30 Phage KC515 DNA was prepared using the procedure described in Genetic Manipulation of *Streptomyces*, A Laboratory Manual, edited by D. Hopwood *et al.* A phage suspension prepared from 10 plates (100 mm) of confluent plaques of KC515 on *S. lividans* TK24 generally gave about 3 µg of phage DNA. The DNA was ligated to circularize at the cos site, subsequently digested with restriction enzymes *Bam*HI and
35 *Pst*I, and dephosphorylated with SAP.

Each module 8 cassette described above was excised with restriction enzymes *Bg*II and *Nsi*I and ligated into the compatible *Bam*HI and *Pst*I sites of KC515 phage DNA prepared as described above. The ligation mixture containing KC515 and various cassettes was transfected into protoplasts of *Streptomyces lividans* TK24 using the
40 procedure described in Genetic Manipulation of *Streptomyces*, A Laboratory Manual edited by D. Hopwood *et al.* and overlaid with TK24 spores. After 16-24 hr, the plaques were restreaked on plates overlaid with TK24 spores. Single plaques were picked and resuspended in 200 µL of nutrient broth. Phage DNA was prepared by the boiling method

- 104 -

(Hopwood *et al.*, *supra*). The PCR with primers spanning the left and right boundaries of the recombinant phage was used to verify the correct phage had been isolated. In most cases, at least 80% of the plaques contained the expected insert. To confirm the presence of the resistance marker (thiostrepton), a spot test is used, as described in Lomovskaya *et al.* (1997), in which a plate with spots of phage is overlaid with mixture of spores of TK24 and phiC31 TK24 lysogen. After overnight incubation, the plate is overlaid with antibiotic in soft agar. A working stock is made of all phage containing desired constructs.

Streptomyces hygroscopicus ATCC 14891 (see US Patent No. 3,244,592, issued 5 Apr 1966, incorporated herein by reference) mycelia were infected with the recombinant phage by mixing the spores and phage (1×10^8 of each), and incubating on R2YE agar (Genetic Manipulation of *Streptomyces*, A Laboratory Manual, edited by D. Hopwood *et al.*) at 30°C for 10 days. Recombinant clones were selected and plated on minimal medium containing thiostrepton (50 µg/ml) to select for the thiostrepton resistance-conferring gene. Primary thiostrepton resistant clones were isolated and purified through a second round of single colony isolation, as necessary. To obtain thiostrepton-sensitive revertants that underwent a second recombination event to evict the phage genome, primary recombinants were propagated in liquid media for two to three days in the absence of thiostrepton and then spread on agar medium without thiostrepton to obtain spores. Spores were plated to obtain about 50 colonies per plate, and thiostrepton sensitive colonies were identified by replica plating onto thiostrepton containing agar medium. The PCR was used to determine which of the thiostrepton sensitive colonies reverted to the wild type (reversal of the initial integration event), and which contain the desired AT swap at module 8 in the ATCC 14891-derived cells. The PCR primers used amplified either the KS/AT junction or the AT/DH junction of the wild-type and the desired recombinant strains. Fermentation of the recombinant strains, followed by isolation of the metabolites and analysis by LCMS, and NMR is used to characterize the novel polyketide compounds.

- 105 -

Example 2

Replacement of Methoxyl with Hydrogen or Methyl at C-13 of FK-506

The present invention also provides the 13-desmethoxy derivatives of FK-506 and the novel PKS enzymes that produce them. A variety of *Streptomyces* strains that produce
5 FK-506 are known in the art, including *S. tsukubaensis* No. 9993 (FERM BP-927), described in U.S. Patent No. 5,624,852, incorporated herein by reference; *S. hygroscopicus* subsp. *yakushimaensis* No. 7238, described in U.S. patent No. 4,894,366, incorporated herein by reference; *S. sp.* MA6858 (ATCC 55098), described in U.S. Patent Nos. 5,116,756, incorporated herein by reference; and *S. sp.* MA 6548, described
10 in Motamedi *et al.*, 1998, "The biosynthetic gene cluster for the macrolactone ring of the immunosuppressant FK-506," *Eur. J. Biochem.* 256: 528-534, and Motamedi *et al.*, 1997, "Structural organization of a multifunctional polyketide synthase involved in the biosynthesis of the macrolide immunosuppressant FK-506," *Eur. J. Biochem.* 244: 74-80, each of which is incorporated herein by reference.

15 The complete sequence of the FK-506 gene cluster from *Streptomyces sp.* MA6548 is known, and the sequences of the corresponding gene clusters from other FK-506-producing organisms is highly homologous thereto. The novel FK-506 recombinant gene clusters of the present invention differ from the naturally occurring gene clusters in that the AT domain of module 8 of the naturally occurring PKSs is replaced by an AT
20 domain specific for malonyl CoA or methylmalonyl CoA. These AT domain replacements are made at the DNA level, following the methodology described in Example 1.

The naturally occurring module 8 sequence for the MA6548 strain is shown below, followed by the illustrative hybrid module 8 sequences for the MA6548 strains.

25 GCATGCGGCTGTACGAGGCGGCACGGCGCACCGGAAGTCCCGTGGTGGTG 50
M R L Y E A A R R T G S P V V V
GCGGCCGCGCTCGACGACGCGCCGGACGTGCCGCTGCTGCGCGGGCTGCG 100
A A A L D D A P D V P L L R G L R
30 GCGTACGACCGTCCGGCGTGCCGCCGTCCGGAACGCTCTCTCGCCGACC 150
R T T V R R A A V R E R S L A D
GCTCGCCGTGCTGCCCGACGACGAGCGCGCCGACGCCTCCCTCGCGTTTCG 200
R S P C C P T T S A P T P P S R S
TCCTGGAACAGCACCGCCACCGTGCTCGGCCACCTGGGCGCCGAAGACAT 250
S W N S T A T V L G H L G A E D I

- 106 -

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CCCCGGCGACGACGACGTTCAAGGAACTCGGCATCGACTCGCTCACC GCGG 300
P A T T T F K E L G I D S L T A
TCCAGCTGCGCAACGCGCTGACCACGGCGACCGCGTACGCCTCAACGCC 350
V Q L R N A L T T A T G V R L N A
ACAGCGGTCTTCGACTTTCGACGCGCGCGCTCGCCGCGAGACTCGG 400
T A V F D F P T P R A L A A R L G
CGACGAGCTGGCCGGTACCCGCGCGCCCGTCGCGGCCCGACCGCGGCCA 450
D E L A G T R A P V A A R T A A
CCGCGGCCCGCGCACGACGAACCGCTGGCGATCGTGGGCATGGCCTGCCGT 500
T A A A H D E P L A I V G M A C R
CTGCCGGGCGGGTTCGCGTCGCCACAGGAGCTGTGGCGTCTCGTCGCGTC 550
L P G G V A S P Q E L W R L V A S
CGGCACCGACGCCATCACGGAGTTCCCCGCGGACCGCGGCTGGGACGTGG 600
G T D A I T E F P A D R G W D V
ACGCGCTCTACGACCCGACCCCGACGCGATCGGCAAGACCTTCGTCCGG 650
D A L Y D P D P D A I G K T F V R
CACGGCGGTTCTTCGACGGTGCGACCGGCTTCGACGCGGCGTTCTTCGG 700
H G G F L D G A T G F D A A F F G
GATCAGCCCGCGCGAGGCCCTGGCCATGGACCCGACGCAACGGGTGCTCC 750
I S P R E A L A M D P Q Q R V L
TGGAGACGTCTTGGGAGGCGTTCGAAAGCGCGGGCATCACCCCGGACGCG 800
L E T S W E A F E S A G I T P D A
GCGCGGGGCGAGCACCCGCGTGTTCATCGGCGCGTTCTCCTACGGGTA 850
A R G S D T G V F I G A F S Y G Y
CGGCAGGGTGCGGATACCAACGGCTTCGGCGCGACAGGGTCGACAGCA 900
G T G A D T N G F G A T G S Q T
GCGTGCTCTCCGGCCGCTCTCGTACTTCTACGGTCTGGAGGGCCCTTCG 950
S V L S G R L S Y F Y G L E G P S
GTCACGGTTCGACACCGCTGCTCGTCGTCCTGGTTCGCCCTGCACCAGGC 1000
V T V D T A C S S S L V A L H Q A
AGGGCAGTCCCTGCGCTCGGGCGAATGCTCGCTCGCCCTGGTCGGCGGTG 1050
G Q S L R S G E C S L A L V G G
TCACGGTGATGGCGTCGCCCCGCGGATTCGTCGAGTTCTCCCGGCGAGCGC 1100
V T V M A S P G G F V E F S R Q R
GGGCTCGCGCCGACGGGCGGGCGAAGGCGTTTCGGCGGGCGCGGACGG 1150
G L A P D G R A K A F G A G A D G
TACGAGCTTCGCCGAGGGCGCCGGTGCCCTGGTGGTCGAGCGGCTCTCCG 1200
T S F A E G A G A L V V E R L S
ACGCGGAGCGCCACGGCCACACCGTCCCTCGCCCTCGTACGCGGCTCCGCG 1250
D A E R H G H T V L A L V R G S A
GCTAACTCCGACGGCGGTCGAACGGTCTGTCGGCGCCGAACGGCCCCCTC 1300
A N S D G A S N G L S A P N G P S
CCAGGAACGCTCATCCACCAGGCCCTCGCGAACGCGAAACTCACCCCG 1350
Q E R V I H Q A L A N A K L T P
CCGATGTCGACGCGGTTCGAGGCGCACGGCACCGGCACCCGCTCGGCGAC 1400
A D V D A V E A H G T G T R L G D
CCCATCGAGGCGCAGGCGCTGCTCGCGACGTACGGACAGGACCGGGCGAC 1450
P I E A Q A L L A T Y G Q D R A T
GCCCCGTGCTGCTCGGCTCGTGAAGTCGAACATCGGGCACGCCCAGGCCG 1500
P L L L G S L K S N I G H A Q A
CGTCAGGGGTGCGCGGATCATCAAGATGGTGCAGGCCATCCGGCACGGG 1550
A S G V A G I I K M V Q A I R H G
GAACTGCCGCCGACACTGCACGCGGACGAGCCGTCGCCGACGTGACTG 1600
E L P P T L H A D E P S P H V D W

- 107 -

5 GACGGCCCGGTGCCGTCGAGCTCCTGACGTCGGCCCCGGCCGTGGCCGGGGA 1650
T A G A V E L L T S A R P W P G
CCGGTCGCCCCGCGCCGCTGCCGTCTCGTCGTTCCGGCGTGAGCGGCACG 1700
T G R P R R A A V S S F G V S G T
10 AACGCCCACATCATCCTTGAGGCAGGACCGGTCAAAACGGGACCGGTCGA 1750
N A H I I L E A G P V K T G P V E
GGCAGGAGCGATCGAGGCAGGACCGGTCTGAAGTAGGACCGGTCTGAGGCTG 1800
A G A I E A G P V E V G P V E A
GACCGCTCCCCGCGGCGCCGCGTCAGCACCGGGCGAAGACCTTCCGCTG 1850
15 G P L P A A P P S A P G E D L P L
CTCGTGTCGGCGCGTTCCCCGGAGGCACTCGACGAGCAGATCGGGCGCCT 1900
L V S A R S P E A L D E Q I G R L
GCGCGCCTATCTCGACACCGGCCCCGGGCGTCGACCGGGCGGCCGTGGCGC 1950
R A Y L D T G P G V D R A A V A
20 AGACACTGGCCCGGCGTACGCACTTCAACCCACCGGGCCGTACTGCTCGGG 2000
Q T L A R R T H F T H R A V L L G
GACACCGTCATCGGCGCTCCCCCGCGGACCAGGCCGACGAACCTCGTCTT 2050
D T V I G A P P A D Q A D E L V F
CGTCTACTCCGGTCAGGGCACCCAGCATCCCGCGATGGGCGAGCAACTCG 2100
25 V Y S G Q G T Q H P A M G E Q L
CGGCCGCGTTCCCCGTGTTCCGCGATGCCTGGCACGACGCGCTCCGACGG 2150
A A A F P V F A D A W H D A L R R
CTCGACGACCCCGACCCGACGACCCACACGGAGCCAGCACACGCTCTT 2200
L D D P D P H D P T R S Q H T L F
30 CGCCCCACAGGCGGCGTTTACCGCCCTCCTGAGGTCCTGGGACATCACGC 2250
A H Q A A F T A L L R S W D I T
CGCACGCCGTCATCGGCCACTCGCTCGGCGAGATCACCGCCGCGTACGCC 2300
P H A V I G H S L G E I T A A Y A
GCCGGGATCCTGTGCTCGACGACGCTGCACCCTGATCACACGCGTGC 2350
35 A G I L S L D D A C T L I T T R A
CCGCCTCATGCACACGCTTCCGCGCGCCGCGCCATGGTCACCGTGCTGA 2400
R L M H T L P P P G A M V T V L
CCAGCGAGGAGGAGGCCCGTCAGGCGCTGCGGCCGGGCGTGGAGATCGCC 2450
T S E E E A R Q A L R P G V E I A
40 GCGGTCTTCCGCCCCGCACTCCGTCTGCTCTCGGGCGACGAGGACGCCGT 2500
A V F G P H S V V L S G D E D A V
GCTCGACGTGCGACAGCGGCTCGGCATCCACCACCGTCTGCCCCGCGCCG 2550
L D V A Q R L G I H H R L P A P
ACGCGGGCCACTCCGCGCACATGGAACCCGTGGCCGCCGAGCTGCTCGCC 2600
45 H A G H S A H M E P V A A E L L A
ACCACTCGCGAGCTCCGTTACGACCGGCCCCACACCGCCATCCCGAACGA 2650
T T R E L R Y D R P H T A I P N D
CCCCACCACCGCCGAGTACTGGGCCGAGCAGGTCCGCAACCCCGTGCTGT 2700
P T T A E Y W A E Q V R N P V L
50 TCCACGCCCACACCCAGCGGTACCCCGACGCCGTGTTCTGTCGAGATCGGC 2750
F H A H T Q R Y P D A V F V E I G
CCCGGCCAGGACCTCTACCGCTGGTCGACGGCATCGCCCTGCAGAACGG 2800
P G Q D L S P L V D G I A L Q N G
CACGGCGGACGAGGTGCACGCGCTGCACACCGCGCTCGCCCGCCTCTTCA 2850
T A D E V H A L H T A L A R L F
CACGCGGCGCCACGCTCGACTGGTCCCGCATCCTCGGCGGTGCTTCGCGG 2900
T R G A T L D W S R I L G G A S R
CACGACCCTGACGTCCCCTCGTACGCGTTCCAGCGGCGTCCCTACTGGAT 2950
H D P D V P S Y A F Q R R P Y W I

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- 108 -

CGAGTCGGCTCCCCCGGCCACGGCCGACTCGGGCCACCCCGTCCTCGGCA 3000
E S A P P A T A D S G H P V L G
CCGGAGTCGCGGTCGCGGGGTCGCGGGGCGGGTGTTCACGGGTCCCGTG 3050
T G V A V A G S P G R V F T G P V
5 CCGCGCGGTGCGGACCGCGCGGTGTTTCATCGCCGAACCTGGCGCTCGCCGC 3100
P A G A D R A V F I A E L A L A A
CGCCGACGCCACCGACTGCGCCACGGTCGAACAGCTCGACGTCACCTCCG 3150
A D A T D C A T V E Q L D V T S
10 TGCCCGGCGGATCCGCGCGGGCAGGGGCCACCGCGCAGACCTGGGTTCGAT 3200
V P G G S A R G R A T A Q T W V D
GAACCCGCGCGGACGGGCGGCGCGCTTCACCGTCCACACCCGCGTCGG 3250
E P A A D G R R R F T V H T R V G
CGACGCCCCGTGGACGCTGCACGCCGAGGGGGTCTCCGCCCCGGCCGCG 3300
D A P W T L H A E G V L R P G R
15 TGCCCCAGCCCGAAGCCGTCGACACCGCCTGGCCCCCGCGGGCGCGGTG 3350
V P Q P E A V D T A W P P P G A V
CCCGCGGACGGGCTGCCCCGGGGCGTGGCGACGCGCGGACCAGGTCTTCGT 3400
P A D G L P G A W R R A D Q V F V
20 CGAAGCCGAAGTCGACAGCCCTGACGGCTTCGTGGCACACCCCGACCTGC 3450
E A E V D S P D G F V A H P D L
TCGACGCGGTCTTCTCCGCGGTGCGCGACGGGAGCCGCCAGCCGACCGGA 3500
L D A V F S A V G D G S R Q P T G
TGGCGCGACCTCGCGGTGCACGCGTCGGACGCCACCGTGCTGCGCGCCTG 3550
W R D L A V H A S D A T V L R A C
25 CCTCACCCGCGCGACAGTGGTGTGCTGGAGCTCGCCGCCTTCGACGGTG 3600
L T R R D S G V V E L A A F D G
CCGGAATGCCGGTGCTCACCGCGGAGTCGGTGACGCTGGGCGAGGTTCGCG 3650
A G M P V L T A E S V T L G E V A
30 TCGGCAGGCGGATCCGACGAGTCGGACGGTCTGCTTCGGCTTGAGTGGTT 3700
S A G G S D E S D G L L R L E W L
GCCGGTGGCGGAGGCCCACTACGACGGTGCCGACGAGCTGCCCGAGGGCT 3750
P V A E A H Y D G A D E L P E G
ACACCCCTCATCACCGCCACACACCCCGACGACCCCGACGACCCACCAAC 3800
Y T L I T A T H P D D P D D P T N
35 CCCCACAACACACCCACACGCACCCACACACAAACCACACGCGTCCTCAC 3850
P H N T P T R T H T Q T T R V L T
CGCCCTCCAACACCACCTCATCACCAACCAACCCCTCATCGTCCACA 3900
A L Q H H L I T T N H T L I V H
40 CCACCACCGACCCCCCAGGCGCGCGCGTCACCGGCCCTACCCGCACCGCA 3950
T T T D P P G A A V T G L T R T A
CAAAACGAACCCCCGCGCATCCACCTCATCGAAACCCACACCCCCA 4000
Q N E H P G R I H L I E T H H P H
CACCCCACTCCCCCTACCCAACCTACCACCCCTCCACCAACCCACCTAC 4050
T P L P L T Q L T T L H Q P H L
45 GCCTCACCAACAACACCCCTCCACACCCCCCACCCTACCCCATCACCAAC 4100
R L T N N T L H T P H L T P I T T
CACCACAACACCACCAACCAACCCCCAACACCCCAACCCCTCAACCCAA 4150
H H N T T T T T P N T P P L N P N
50 CCACGCCATCCTCATCACCGGCGGCTCCGGCACCCCTCGCCGGCATCCTCG 4200
H A I L I T G G S G T L A G I L
CCCGCCACCTCAACCAACCCCAACCTACCTCCTCTCCCGCACACCACCA 4250
A R H L N H P H T Y L L S R T P P
CCCCCAACCAACCCGGCACCCACATCCCTGCGACCTACCGACCCAC 4300
P P T T P G T H I P C D L T D P T

- 109 -

CCAAATCACCCAAGCCCTCACCCACATACCACAACCCCTCACCGGCATCT 4350
Q I T Q A L T H I P Q P L T G I
TCCACACCGCCGCCACCCCTCGACGACGCCACCCCTCACCAACCTCACCCCC 4400
F H T A A T L D D A T L T N L T P
5 CAACACCTCACCCACCCCTCCAACCCAAAGCCGACGCCGCTGGCACCT 4450
Q H L T T T L Q P K A D A A W H L
CCACCACCACACCCAAAACCAACCCCTCACCCACTTCGTCCTCTACTCCA 4500
H H H T Q N Q P L T H F V L Y S
GCGCGCGCCGCCACCCCTCGGCAGCCCCGGCCAAGCCAACTACGCCGCGGCC 4550
10 S A A A T L G S P G Q A N Y A A A
AACGCCTTCTCGACGCCCTCGCCACCCACCGCCACACCCAAGGACAACC 4600
N A F L D A L A T H R H T Q G Q P
CGCCACCACCATCGCCTGGGGCATGTGGCACACCACCACACTCACCA 4650
A T T I A W G M W H T T T T L T
15 GCCAACTCACCGACAGCGACCGCGACCGCATCCGCCGCGGCGGCTTCTTG 4700
S Q L T D S D R D R I R R G G F L
CCGATCTCGGACGACGAGGGCATGC
P I S D D E G M

20 The *AvrII-XhoI* hybrid FK-506 PKS module 8 containing the AT domain of
module 12 of rapamycin is shown below.

GCATGCGGCTGTACGAGGCGGCACGGCGCACCGGAAGTCCCGTGGTGGTG 50
M R L Y E A A R R T G S P V V V
GCGGCCGCGCTCGACGACGCGCGGACGTGCCGCTGCTGCGCGGGCTGCG 100
25 A A A L D D A P D V P L L R G L R
GCGTACGACCGTCCGGCGTGCCGCCGTCCGGGAACGCTCTCTCGCCGACC 150
R T T V R R A A V R E R S L A D
GCTCGCCGTGCTGCCCCGACGACGAGCGCGCCGACGCCTCCCTCGCGTTG 200
R S P C C P T T S A P T P P S R S
30 TCCTGGAACAGCACCGCCACCGTGCTCGGCCACCTGGGCGCCGAAGACAT 250
S W N S T A T V L G H L G A E D I
CCCGGCGACGACGACGTTCAAGGAACCTCGGCATCGACTCGCTCACCGCGG 300
P A T T T F K E L G I D S L T A
TCCAGTGTGCGCAACGCGCTGACCACGGCGACCGGCTACGCCTCAACGCC 350
35 V Q L R N A L T T A T G V R L N A
ACAGCGGTCTTCGACTTTCCGACGCGCGCGCTCGCCGCGAGACTCGG 400
T A V F D F P T P R A L A A R L G
CGACGAGTGGCCGGTACCCGCGCGCCCGTCCGCGCCCGGACCGCGGCCA 450
D E L A G T R A P V A A R T A A
40 CCGCGGCCGCGCACGACGAACCGCTGGCGATCGTGGGCATGGCCTGCCGT 500
T A A A H D E P L A I V G M A C R
CTGCCGGGCGGGGCTCGCGTCGCCACAGGAGCTGTGGCGTCTCGTCGCGTC 550
L P G G V A S P Q E L W R L V A S
CGGCACCGACGCCATCACGGAGTTCCCCGCGGACCGCGGCTGGGACGTGG 600
45 G T D A I T E F P A D R G W D V
ACGCGCTCTACGACCCGGACCCCGACGCGATCGGCAAGACCTTCGTCCGG 650
D A L Y D P D P D A I G K T F V R
CACGGCGGCTTCTCGACGGTGCGACCGGCTTCGACGCGGCGTTCTTCGG 700
H G G F L D G A T G F D A A F F G
50 GATCAGCCCGCGGAGGCCCTGGCCATGGACCCGAGCAACGGGTGCTCC 750
I S P R E A L A M D P Q Q R V L
TGGAGACGTCTGGGAGGCGTTCGAAAGCGCGGCATACCCCGGACGCG 800

- 110 -

L E T S W E A F E S A G I T P D A
GCGCGGGGACGACACCGGCGTGTTCATCGGCGCGTTCTCCTACGGGTA 850
A R G S D T G V F I G A F S Y G Y
CGGCACGGGTGCGGATACCAACGGCTTCGGCGCGACAGGGTCGCAGACCA 900
5 G T G A D T N G F G A T G S Q T
GCGTGCTCTCCGGCCGCTCTCGTACTTCTACGGTCTGGAGGGCCCTTCG 950
S V L S G R L S Y F Y G L E G P S
GTCACGGTCGACACCGCCTGCTCGTCGTCGTCGCTGCGCCCTGCACCAGGC 1000
V T V D T A C S S S L V A L H Q A
10 AGGGCAGTCCCTGCGCTCGGGCGAATGCTCGCTCGCCCTGGTCGGCGGTG 1050
G Q S L R S G E C S L A L V G G
TCACGGTGATGGCGTCGCGCGGCGGATTCTCGAGTTCTCCCGGCAGCGC 1100
V T V M A S P G G F V E F S R Q R
GGGCTCGCGCCGGACGGGCGGGCGAAGGCGTTTCGGCGCGGGCGCGGACGG 1150
15 G L A P D G R A K A F G A G A D G
TACGAGCTTCGCCGAGGGCGCCGGTGCCCTGGTGGTCGAGCGGCTCTCCG 1200
T S F A E G A G A L V V E R L S
ACGCGGAGCGCCACGGCCACACCGTCTCGCCCTCGTACGCGGCTCCGCG 1250
D A E R H G H T V L A L V R G S A
20 GCTAATCCGACGGCGCGTCGAACGGTCTGTCGGCGCCGAACGGCCCCCTC 1300
A N S D G A S N G L S A P N G P S
CCAGGAACGCGTCATCCACCAGGCCCTCGCGAACGCGAAACTCACCCCG 1350
Q E R V I H Q A L A N A K L T P
CCGATGTCGACGCGGTTCGAGGCGCACGGCACCGGCACCCGCTCGGCGAC 1400
25 A D V D A V E A H G T G T R L G D
CCCATCGAGGCGCAGGCGCTGCTCGCGACGTACGGACAGGACCGGGCGAC 1450
P I E A Q A L L A T Y G Q D R A T
GCCCCTGCTGCTCGGCTCGCTGAAGTCGAACATCGGGCACGCCAGGCCG 1500
P L L L G S L K S N I G H A Q A
30 CGTCAGGGGTGCGCGGGATCATCAAGATGGTGCAGGCCATCCGGCACGGG 1550
A S G V A G I I K M V Q A I R H G
GAACTGCCGCCGACACTGCACGCGGACGAGCCGTCGCCGCACGTCGACTG 1600
E L P P T L H A D E P S P H V D W
GACGGCCCGGTGCCGTGAGCTCCTGACGTGCGGCCCGGCCGTGGCCGGGGA 1650
35 T A G A V E L L T S A R P W P G
CCGGTCGCCCCTAGGCGGGCAGGCGTGTCTGCTCCTTCGGGATCAGTGGCACC 1700
T G R P R R A G V S S F G I S G T
AACGCCACGTATCTGGAAGCGCACCCCCCACTCAGCCTGCGGACAA 1750
N A H V I L E S A P P T Q P A D N
40 CGCGGTGATCGAGCGGCACCGGAGTGGGTGCCGTTGGTGATTTTCGGCCA 1800
A V I E R A P E W V P L V I S A
GGACCCAGTCGGCTTTGACTGAGCACGAGGGCCGGTTGCGTGCGTATCTG 1850
R T Q S A L T E H E G R L R A Y L
GCGGCGTCGCCCCGGGTGGATATGCGGGCTGTGGCATCGACGCTGGCGAT 1900
45 A A S P G V D M R A V A S T L A M
GACACGGTCGGTGTTTCGAGCACCGTGCCGTGCTGCTGGGAGATGACACCG 1950
T R S V F E H R A V L L G D D T
TCACCGGCACCGCTGTGTCTGACCCTCGGGCGGTGTTCTGCTTTCCCGGGA 2000
V T G T A V S D P R A V F V F P G
50 CAGGGGTGCGCAGCGTGTGGCATGGGTGAGGAACTGGCCGCCGCGTTCCC 2050
Q G S Q R A G M G E E L A A A F P
CGTCTTCGCGCGGATCCATCAGCAGGTGTGGGACCTGCTCGATGTGCCCG 2100
V F A R I H Q Q V W D L L D V P
ATCTGGAGGTGAACGAGACCGGTTACGCCCAGCCGGCCCTGTTTCGCAATG 2150

SEQ ID NO: 1

- 111 -

D L E V N E T G Y A Q P A L F A M
CAGGTGGCTCTGTTCCGGGCTGCTGGAATCGTGGGGTGTACGACCGGACGC 2200
Q V A L F G L L E S W G V R P D A
GGTGATCGGCCATTCCGGTGGGTGAGCTTGC GGCTGCGTATGTGTCCGGGG 2250
5 V I G H S V G E L A A A Y V S G
TGTGGTTCGTGGAGGATGCCTGCACTTTGGTGTCCGGCGGGGCTCGTCTG 2300
V W S L E D A C T L V S A R A R L
ATGCAGGCTCTGCCCCGGGTGGGGTGTGGTGTGTCCCGGTCTCGGA 2350
M Q A L P A G G V M V A V P V S E
10 GGATGAGGCCCGGGCCGTGCTGGGTGAGGGTGTGGAGATCGCCGCGGTCA 2400
D E A R A V L G E G V E I A A V
ACGGCCCCGTGCTCGGTGGTTCTCTCCGGTGTGAGGCCCGGTGCTGCAG 2450
N G P S S V V L S G D E A A V L Q
GCCGCGGAGGGGCTGGGGAAGTGGACGCGGTGGCGACCAGCCACGCGTT 2500
15 A A E G L G K W T R L A T S H A F
CCATTCCGCCCGTATGGAACCCATGCTGGAGGAGTTCCGGGCGGTGCGCCG 2550
H S A R M E P M L E E F R A V A
AAGGCCTGACCTACCGGACGCCGAGGTCTCCATGGCCGTTGGTGATCAG 2600
E G L T Y R T P Q V S M A V G D Q
20 GTGACCACCGCTGAGTACTGGGTGCGGCAGGTCCGGGACACGGTCCGGTT 2650
V T T A E Y W V R Q V R D T V R F
CGGCGAGCAGGTGGCCTCGTACGAGGACGCCGTGTTGTCGTCGAGCTGGGTG 2700
G E Q V A S Y E D A V F V E L G
CCGACCGGTCACTGGCCCGCTGGTTCGACGGTGTCCGCGATGCTGCACGGC 2750
25 A D R S L A R L V D G V A M L H G
GACCACGAAATCCAGGCCGCGATCGGCGCCCTGGCCACCTGTATGTCAA 2800
D H E I Q A A I G A L A H L Y V N
CGGCGTCACGGTCACTGGCCCGCGCTCCTGGGCGATGCTCCGGCAACAC 2850
G V T V D W P A L L G D A P A T
30 GGGTGTGACCTTCCGACATACGCTTCCAGCACCAGCGCTACTGGCTC 2900
R V L D L P T Y A F Q H Q R Y W L
GAGTCGGCTCCCCCGGCCACGGCCGACTCGGGCCACCCCGTCTCCGGCAC 2950
E S A P P A T A D S G H P V L G T
CGGAGTCGCCGTCGCCGGGTGCCGGGGCGGGTGTTCACGGGTCCCGTGC 3000
35 G V A V A G S P G R V F T G P V
CCGCCGGTGCAGCCGCGCGGTGTTTCATCGCCGAACGGCGCTCGCCGCC 3050
P A G A D R A V F I A E L A L A A
GCCGACGCCACCGACTGCGCCACGGTCAACAGCTCGACGTCACCTCCGT 3100
A D A T D C A T V E Q L D V T S V
40 GCGCGGCGGATCCGCGCGCGGCAGGGCCACCGCGCAGACCTGGGTGATG 3150
P G G S A R G R A T A Q T W V D
AACCCGCCGCCGACGGGCGGCGCGCTTACCGTCCACACCCGCGTCCGGC 3200
E P A A D G R R R F T V H T R V G
GACGCCCCGTGGACGCTGCACGCCGAGGGGGTTCTCCGCCCCGGCGCGT 3250
45 D A P W T L H A E G V L R P G R V
GCCCCAGCCCGAAGCCGTGACACCGCCTGGCCCCCGCGGGCGCGGTGC 3300
P Q P E A V D T A W P P P G A V
CCGCGGACGGGCTGCCCCGGGCGTGGCGACGCGCGGACCAGGTCTTCGTC 3350
P A D G L P G A W R R A D Q V F V
50 GAAGCCGAAGTCGACAGCCCTGACGGCTTCGTGGCACACCCCGACCTGCT 3400
E A E V D S P D G F V A H P D L L
CGACGCGGTCTTCTCCGCGGTGCGCGACGGGAGCCGCCAGCCGACCGGAT 3450
D A V F S A V G D G S R Q P T G
GGCGGACCTCGCGGTGCACGCGTCCGACGCCACCGTGTGCGCGCTGC 3500

- 112 -

W R D L A V H A S D A T V L R A C
CTCACCCGCCGCGACAGTGGTGTCTGGAGCTCGCCGCCTTCGACGGTGC 3550
L T R R D S G V V E L A A F D G A
CGGAATGCCGGTGCTCACCGCGGAGTCGGTGACGCTGGGCGAGGTGCGGT 3600
5 G M P V L T A E S V T L G E V A
CGGCAGGCGGATCCGACGAGTCGGACGGTCTGCTTCGGCTTGAGTGGTTG 3650
S A G G S D E S D G L L R L E W L
CCGGTGGCGGAGGCCACTACGACGGTGCCGACGAGCTGCCCAGGGGCTA 3700
P V A E A H Y D G A D E L P E G Y
10 CACCCTCATCACCGCCACACACCCCGACGACCCCGACGACCCACCAACC 3750
T L I T A T H P D D P D D P T N
CCCACAACACACCCACACGCACCCACACACAAACCACACGCGTCCTCACC 3800
P H N T P T R T H T Q T T R V L T
GCCCTCCAACACCACCTCATCACCACCAACCACACCCCTCATCGTCCACAC 3850
15 A L Q H H L I T T N H T L I V H T
CACCACCGACCCCCAGGCGCCGCGTACCAGGCTCACCAGCACCAC 3900
T T D P P G A A V T G L T R T A
AAAACGAACACCCCGGCGCATCCACCTCATCGAAACCCACCACCCCCAC 3950
Q N E H P G R I H L I E T H H P H
20 ACCCACTCCCCCTCACCAACTCACCACCCTCCACCAACCCACCTACG 4000
T P L P L T Q L T T L H Q P H L R
CCTCACCAACACCCCTCCACACCCCCACCTCACCCTCATCACCACCC 4050
L T N N T L H T P H L T P I T T
ACCACAACACCACCAACACCCCAACACCCACCCCTCAACCCCAAC 4100
25 H H N T T T T T P N T P P L N P N
CAGCCATCCTCATCACCGGCGGCTCCGGCACCCCTCGCCGGCATCCTCGC 4150
H A I L I T G G S G T L A G I L A
CCGCCACCTCAACACCCCCACACCTACCTCCTCTCCGCACACCACCAC 4200
R H L N H P H T Y L L S R T P P
30 CCCCCACACACCCGGCACCCACATCCCCTGCGACCTCACCAGCCCCACC 4250
P P T T P G T H I P C D L T D P T
CAAATACCCAAGCCCTCACCACATAACCACAACCCCTCACCAGCATCTT 4300
Q I T Q A L T H I P Q P L T G I F
CCACACCGCCGCCACCTCGACGACGCCACCCCTACCAACCTCACCCTCC 4350
35 H T A A T L D D A T L T N L T P
AACACCTCACCACCACCTCCAACCCAAAGCCGACGCCGCTGGCACCTC 4400
Q H L T T T L Q P K A D A A W H L
CACCACCACACCCAAAACCAACCCCTCACCCTTCTGTCCTCTACTCCAG 4450
H H H T Q N Q P L T H F V L Y S S
40 CGCCGCCGCCACCTCGGCAGCCCCGGCCAAGCCAACCTACGCCGCCGCCA 4500
A A A T L G S P G Q A N Y A A A
ACGCCTTCTCGACGCCCTCGCCACCCACCGCCACACCAAGGACAACCC 4550
N A F L D A L A T H R H T Q G Q P
GCCACCACCATCGCCTGGGGCATGTGGCACACCACCACCACTCACCAG 4600
45 A T T I A W G M W H T T T T L T S
CCAACCTACCGACAGCGACCGCGACCGCATCCGCCGCGGCGGCTTCCTGC 4650
Q L T D S D R D R I R R G G F L
CGATCTCGGACGACGAGGGCATGC
P I S D D E G M
50

The *AvrII-XhoI* hybrid FK-506 PKS module 8 containing the AT domain of module 13 of rapamycin is shown below.

Figure 6

Figure 6 displays two sets of plots comparing the performance of different models across various metrics. The top set of plots shows the results for the "Model A" series, while the bottom set shows the results for the "Model B" series. Each plot includes a legend indicating the different components being compared.

The top set of plots (Model A) includes:

- A bar chart showing the distribution of values across categories.
- A line graph showing the trend over time or iterations.
- A scatter plot showing the relationship between variables.

The bottom set of plots (Model B) includes:

- A bar chart showing the distribution of values across categories.
- A line graph showing the trend over time or iterations.
- A scatter plot showing the relationship between variables.

The plots illustrate how the models perform under different conditions, highlighting the strengths and weaknesses of each approach.

[illegible]

- 114 -

CCGATGTCGACGCGGTCGAGGCGCACGGCACCCGCGCTCGGCGAC 1400
A D V D A V E A H G T G T R L G D
CCCATCGAGGCGCAGGCGCTGCTCGCGACGTACGGACAGGACCGGGCGAC 1450
P I E A Q A L L A T Y G Q D R A T
5 GCCCCTGCTGCTCGGCTCGCTGAAGTCGAACATCGGGCACGCCAGGCCG 1500
P L L L G S L K S N I G H A Q A
CGTCAGGGGTCGCCGGGATCATCAAGATGGTGCAGGCCATCCGGCACGGG 1550
A S G V A G I I K M V Q A I R H G
10 GAACTGCCGCCGACACTGCACGCGGACGAGCCGTGCGCGACGTGACTG 1600
E L P P T L H A D E P S P H V D W
GACGGCCGGTGCCGTCGAGCTCCTGACGTGCGCCCCGGCCGTGGCCGGGGA 1650
T A G A V E L L T S A R P W P G
CCGGTCGCCCTAGGCGGGCGGGCGTGTCTCCTTCGGAGTCAGCGGCACC 1700
T G R P R R A G V S S F G V S G T
15 AACGCCCACGTCATCCTGGAGAGCGCACCCCCGCTCAGCCCGCGGAGGA 1750
N A H V I L E S A P P A Q P A E E
GGCGCAGCCTGTTGAGACGCCGGTGGTGGCCTCGGATGTGCTGCCGCTGG 1800
A Q P V E T P V V A S D V L P L
TGATATCGGCCAAGACCCAGCCCGCCCTGACCGAACACGAAGACCGGCTG 1850
20 V I S A K T Q P A L T E H E D R L
CGCGCCTACCTGGCGGCGTCGCCCGGGGCGGATATACGGGCTGTGGCATC 1900
R A Y L A A S P G A D I R A V A S
GACGCTGGCGGTGACACGGTCGGTGTTCGAGCACCGCGCCGTACTCCTTG 1950
T L A V T R S V F E H R A V L L
25 GAGATGACACCGTCACCGGCACCGCGGTGACCGACCCAGGATCGTGTTT 2000
G D D T V T G T A V T D P R I V F
GTCTTTCCCGGGCAGGGGTGGCAGTGGCTGGGGATGGGCAGTGCAGTGC 2050
V F P G Q G W Q W L G M G S A L R
CGATTGCTCGGTGGTGTTCGCCGAGCGGATGGCCGAGTGTGCGGCGGCGT 2100
30 D S S V V F A E R M A E C A A A
TGCGCGAGTTCTGTTGACTGGGATCTGTTACGGTTCTGGATGATCCGGCG 2150
L R E F V D W D L F T V L D D P A
GTGGTGGACCGGGTTGATGTGGTCCAGCCCGCTTCCTGGGCGATGATGGT 2200
V V D R V D V V Q P A S W A M M V
35 TTCCCTGGCCGCGGTGTGGCAGGCGGCCGGTGTGCGGCCGGATGCGGTGA 2250
S L A A V W Q A A G V R P D A V
TCGGCCATTTCGAGGGTGAAGATCGCCGAGCTTGTGTGGCGGGTGGCGTG 2300
I G H S Q G E I A A A C V A G A V
TCACTACGCGATGCCGCCCGGATCGTGACCTTGCGCAGCCAGGCGATCGC 2350
40 S L R D A A R I V T L R S Q A I A
CCGGGGCCTGGCGGGGCCGGGCGCGATGGCATCCGTCGCCCTGCCCGCGC 2400
R G L A G R G A M A S V A L P A
AGGATGTGAGCTGGTCGACGGGGCCTGGATCGCCGCCACAACGGGCCC 2450
Q D V E L V D G A W I A A H N G P
45 GCCTCCACCGTGATCGCGGGCACCCCGGAAGCGGTGACCATGTCCTCAC 2500
A S T V I A G T P E A V D H V L T
CGCTCATGAGGCACAAGGGGTGCGGGTGGCGCGGATCACCGTCGACTATG 2550
A H E A Q G V R V R R I T V D Y
CCTCGCACACCCCGCACGTCGAGCTGATCCGCGACGAAGTACTCGACATC 2600
50 A S H T P H V E L I R D E L L D I
ACTAGCGACAGCAGCTCGCAGACCCCGCTCGTGCCGTGGCTGTGACCGT 2650
T S D S S S Q T P L V P W L S T V
GGACGGCACCTGGGTGACAGCCCGCTGGACGGGGAGTACTGGTACCGGA 2700
D G T W V D S P L D G E Y W Y R

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ACCTGCGTGAACCTGGGTCTGGTTTCCACCCCGCCGCTGAGCCAGTTGCAGGCC 2750
N L R E P V G F H P A V S Q L Q A
CAGGGCGACACCGTGTTCGTCGAGGTACAGCGCCAGCCCGGTGTTGTTGCA 2800
Q G D T V F V E V S A S P V L L Q
GGCGATGGACGACGATGTGCTCACGGTTGCCACGCTGCGTCGTGACGACG 2850
A M D D D V V T V A T L R R D D
GCGACGCCACCCGGATGCTCACCGCCCTGGCACAGGCCTATGTCCACGGC 2900
G D A T R M L T A L A Q A Y V H G
GTCACCGTCGACTGGCCCGCCATCCTCGGCACCACCACAACCCGGGTACT 2950
V T V D W P A I L G T T T T R V L
GGACCTTCCGACCTACGCCTTCCAACACCAGCGGTACTGGCTCGAGTCGG 3000
D L C P T Y A F Q Q H Q R Y W L E S
CTCCCCCGGCCACGGCCGACTCGGGCCACCCCGTCTCGGCACCGGAGTC 3050
A P P A T A D S G H P V L G T G G V
GCCGTGCGCCGGGTGCGCCGGGCCGGGTGTTACGGGTCCCGTGCCCGCCGG 3100
A V A G S P G R V F T G P V P A G
TGCGGACCGCGCGGTGTTTCATCGCCGAAGTGGCGCTCGCCGCCGCCGACG 3150
A D R A V F I A E L A L A A A D
CCACCGACTGCGCCACGGTCTGAACAGCTCGACGTCACCTCCGTGCCCGGC 3200
A T D C A T V E Q L D V T S V P G
GGATCCGCCCCGCGGCAGGGCCACCGCGCAGACCTGGGTGCGATGAACCCGC 3250
G S A R G R A T A Q T W V D E P A
CGCCGACGGGCGGCGCGGCTTACCGTCCACACCCGCGTCGGCGACGCCC 3300
A D G R R R F T V H T R V G D A
CGTGGACGCTGCACGCCGAGGGGGTTCTCCGCCCCGCGCGCTGCCCCAG 3350
P W T L H A E G V L R P G R V P Q
CCCGAAGCCGTCGACACCGCCTGGCCCCCGCCGGGCGCGGTGCCCGCGGA 3400
P E A V D T A W P P P G A V P A D
CGGGCTGCCCGGGGCGTGGCGACGCGCGGACCAGGTCTTCGTCTGAAGCCG 3450
G L P G A W R R A D Q V F V E A
AAGTCGACAGCCCTGACGGCTTCGTGGCACACCCCGACCTGCTCGACGCG 3500
E V D S P D G G F V A H P D L D G A
GTCTTCTCCGCGGTGCGGACGCGGAGCCGACCGACCGGATGGCGGA 3550
V F S A V G D G S R Q P T G W R D
CCTCGCGGTGCACGCGTCGGACGCCACCGTGCTGCGCGCCTGCCTCACCC 3600
L A V H A S D A T V L R A C L T
GCCGCGACAGTGGTGTGCTGGAGCTCGCCGCCTTCGACGGTGCCGGAATG 3650
R R D S G V V E L A A F D G A G M
CCGGTGCTCACCGCGGAGTCGGTGACGCTGGGCGAGGTGCGCTCGGCAGG 3700
P V L T A E S V T L G E V A S A G
CGGATCCGACGAGTCGGACGGTCTGCTTCGGCTTGAGTGTTGCCGGTGG 3750
G S D E S D G L L R L E W L P V
CGGAGGCCCACTACGACGGTGCCGACGAGCTGCCCGAGGGCTACACCCTC 3800
A E A H Y D G A D E L P E G Y T L
ATCACCGCCACACACCCCGACGACCCCGACGACCCACCAACCCCCACAA 3850
I T A T H P D D P D D P T N P H N
CACACCCACACGCACCCACACACAAACCACACGCGTCCTCACCGCCCTCC 3900
T P T R T H T Q T T R V L T A L
AACACCCTCATCACCACCAACCACACCTCATCGTCCACACCACCACC 3950
Q H H L I C T T N H T L I V H T T T
GACCCCCCAGGCGCGCGTACCGGCCTACCCGACCCGCACAAAACGA 4000
D P P G A A V T G L T R T A Q N E
ACACCCCGCGCGCATCCACCTCATCGAAACCCACCACCCCCACACCCAC 4050
H P G R I H L I E T H H P H T P

- 116 -

TCCCCCTACCCAACTCACCACCCTCCACCAACCCACCTACGCCTCACC 4100
 L P L T Q L T T L H Q P H L R L T
 AACAAACACCCTCCACACCCCCACCTCACCCTCATCACCACCCACCACAA 4150
 N N T L H T P H L T P I T T H H N
 5 CACCACCACAACCACCCCAACACCCACCCCTCAACCCCAACCACGCCA 4200
 T T T T T P N T P P L N P N H A
 TCCTCATCACCAGCGGCTCCGGCACCCCTCGCCGGCATCCTCGCCCGCCAC 4250
 I L I T G G S G T L A G I L A R H
 CTCAACCACCCCAACACCTACCTCCTCTCCCGCACACCACCCACCCAC 4300
 10 L N H P H T Y L L S R T P P P P T
 CACACCCGGCACCCACATCCCCTGCGACCTCACCACCCCAACCAATCA 4350
 T P G T H I P C D L T D P T Q I
 CCAAGCCCTCACCACATACCACAACCCCTCACCAGCATCTTCCACACC 4400
 T Q A L T H I P Q P L T G I F H T
 15 GCCGCCACCCTCGACGACGCCACCCTCACCACCTCACCACCCCAACACCT 4450
 A A T L D D A T L T N L T P Q H L
 CACCACCACCCTCCAACCCAAAGCCGACGCCGCTGGCACCTCCACCACC 4500
 T T T L Q P K A D A A W H L H H
 ACACCCAAAACCAACCCCTCACCACCTTCGTCCTCTACTCCAGCGCCGCC 4550
 20 H T Q N Q P L T H F V L Y S S A A
 GCCACCTCGGCACCCCGCAAGCCAACTACGCCGCGGCCAACGCCTT 4600
 A T L G S P G Q A N Y A A A N A F
 CCTCGACGCCCTCGCCACCCACCGCCACACCAAGGACAACCCGCCACCA 4600
 L D A L A T H R H T Q G Q P A T
 25 CCATCGCCTGGGGCATGTGGCACACCACCACACTCACCAGCCAACTC 4700
 T I A W G M W H T T T T L T S Q L
 ACCGACAGCGACCGCGACCGCATCCGCCGCGCGGCTTCTGCCGATCTC 4750
 T D S D R D R I R R G G F L P I S
 GGACGACGAGGGCATGC
 30 D D E G M

The *NheI-XhoI* hybrid FK-506 PKS module 8 containing the AT domain of module 12 of rapamycin is shown below.

GCATGCGGCTGTACGAGGCGGCACGGCGCACCGGAAGTCCCGTGGTGGTG 50
 35 M R L Y E A A R R T G S P V V V
 GCGGCGCGCTCGACGACGCGCGGACGTGCCGCTGCTGCGCGGGCTGCG 100
 A A A L D D A P D V P L L R G L R
 GCGTACGACCGTCCGGCGTCCGCGCTCCGGGAACGCTCTCTCGCCGACC 150
 R T T V R R A A V R E R S L A D
 40 GCTCGCCGTGCTGCCCCGACGACGAGCGCGCGGACGCCTCCCTCGCGTTCTG 200
 R S P C C P T T S A P T P P S R S
 TCCTGGAACAGCACCGCCACCGTGCTCGGCCACCTGGGCGCCGAAGACAT 250
 S W N S T A T V L G H L G A E D I
 CCCGGCGACGACGACGTTCAAGGAACCTCGGCATCGACTCGCTCACCAGCGG 300
 45 P A T T T F K E L G I D S L T A
 TCCAGCTGCGCAACGCGCTGACCACGGCGACCGGCGTACGCCTCAACGCC 350
 V Q L R N A L T T A T G V R L N A
 ACAGCGGTCTTCGACTTTCGACGCGCGCGGCTCGCCGCGAGACTCGG 400
 T A V F F P T P R A L A A R L G
 50 CGACGAGCTGGCCGTTACCGCGCGCGCGTCCGCGGCCGACCGCGGCCA 450
 D E L A G T R A P V A A R T A A
 CCGCGCGCGCGCACGACGAACCGCTGGCGATCGTGGGCATGGCCTGCCGT 500

- 117 -

Sequence

T A A A H D E P L A I V G M A C R
CTGCCGGGCGGGGTGCGCTCGCCACAGGAGCTGTGGCGTCTCGTCGCGTC 550
L P G G V A S P Q E L W R L V A S
CGGCACCGACGCCATCACGGAGTTCCCCGCGGACCGCGGCTGGGACGTGG 600
5 G T D A I T E F P A D R G W D V
ACGCGCTCTACGACCCGGACCCCGACGCGATCGGCAAGACCTTCGTCCGG 650
D A L Y D P D P D A I G K T F V R
CACGGCGGCTTCTCGACGGTGCAGCCGGCTTCGACGCGGCGTTCCTTCGG 700
H G G F L D G A T G F D A A F F G
10 GATCAGCCCGCGCGAGGCCCTGGCCATGGACCCGAGCAACGGGTGCTCC 750
I S P R E A L A M D P Q Q R V L
TGGAGACGTCCTGGGAGGCGTTCGAAAGCGCGGGCATCACCCCGGACGCG 800
L E T S W E A F E S A G I T P D A
GCGCGGGGCGAGCGACACCGGCGTGTTCATCGGCGGCTTCTCCTACGGGTA 850
15 A R G S D T G V F I G A F S Y G Y
CGGCACGGGTGCGGATACCAACGGCTTCGGCGCGACAGGGTCGCAGACCA 900
G T G A D T N G F G A T G S Q T
GCGTGCTCTCCGGCCGCTCTCGTACTTCTACGGTCTGGAGGGCCCTTCG 950
S V L S G R L S Y F Y G L E G P S
20 GTCACGGTCGACACCGCTCGTCTCGTCACTGGTCGCCCTGCACCAGGC 1000
V T V D T A C S S S L V A L H Q A
AGGGCAGTCCCTGCGCTCGGGCGAATGCTCGCTCGCCCTGGTCGGCGGTG 1050
G Q S L R S G E C S L A L V G G
TCACGGTGATGGCGTCGCCCGGCGGATTCTGTCGAGTTCTCCCGGCAGCGC 1100
25 V T V M A S P G G F V E F S R Q R
GGGCTCGCGCCGACGGGCGGGCGAAGGCGTTCGGCGCGGGCGCGGACGG 1150
G L A P D G R A K A F G A G A D G
TACGAGCTTCGCCGAGGGCGCCGGTGCCTGGTGGTCGAGCGGCTCTCCG 1200
T S F A E G A G A L V V E R L S
30 ACGCGGAGCGCCACGGCCACACCGTCTCGCCCTCGTACGCGGCTCCGCG 1250
D A E R H G H T V L A L V R G S A
GCTAACTCCGACGGCGCGTCAACGGTCTGTGCGGCGCCGAACGGCCCCCTC 1300
A N S D G A S N G L S A P N G P S
CCAGGAACGCGTCATCCACCAGGCCCTCGCGAACGCGAAACTCACCCCG 1350
35 Q E R V I H Q A L A N A K L T P
CCGATGTGACGCGGTGAGGCGCACGGCACCGGCACCCGCTCGGCGAC 1400
A D V D A V E A H G T G T R L G D
CCCATCGAGGCGCAGGCGTGTCTCGCGACGTACGGACAGGACCGGGCGAC 1450
P I E A Q A L L A T Y G Q D R A T
40 GCCCCTGTGCTCGGCTCGCTGAAGTCGAACATCGGGCACGCCCAGGCCG 1500
P L L L G S L K S N I G H A Q A
CGTCAGGGGTGCGCGGGATCATCAAGATGGTGCAGGCCATCCGGCACGGG 1550
A S G V A G I I K M V Q A I R H G
GAACTGCCGCCGACACTGCACGCGGACGAGCCGTGCGCGCACGTGCGACTG 1600
45 E L P P T L H A D E P S P H V D W
GACGGCCGGTGCCGTGAGCTCCTGACGTGCGGCCGCGGCTGGCCGGGGA 1650
T A G A V E L L T S A R P W P G
CCGGTCGCCCCGCGCCGCGCTGCCGTCTCGTCGTTTCGGCGTGAGCGGCACG 1700
T G R P R R A A V S S F G V S G T
50 AACGCCCACATCATCTTGAGGCAGGACCGGTCAAAACGGGACCGGTGCA 1750
N A H I I L E A G P V K T G P V E
GGCAGGAGCGATCGAGGCAGGACCGGTGCAAGTAGGACCGGTGAGGCTG 1800
A G A I E A G P V E V G P V E A
GACCGCTCCCCGCGGCGCCGCGTCAACCGGGCGAAGACCTTCCGCTG 1850

- 118 -

G P L P A A P P S A P G E D L P L
CTCGTGTGCGGCGGTTCCCCGGAGGCACTCGACGAGCAGATCGGGCGCCT 1900
L V S A R S P E A L D E Q I G R L
GCGGCGCTATCTCGACACCGGCCCCGGGCGTTCGACCGGGCGGCGCTGGCGC 1950
5 R A Y L D T G P G V D R A A V A
AGACACTGGCCCGGCGTACGCACTTCACCCACCGGGCGTACTGCTCGGG 2000
Q T L A R R T H F T H R A V L L G
GACACCGTCATCGGCGCTCCCCCGCGGACCAGGCCGACGAACCTCGTCTT 2050
D T V I G A P P A D Q A D E L V F
10 CGTCTACTCCGGTCAGGGCACCCAGCATCCCGCGATGGGCGAGCAGCTAG 2100
V Y S G Q G T Q H P A M G E Q L
CCGCCGCGTTCCCCGTTCTCGCGCGGATCCATCAGCAGGTGTGGGACCTG 2150
A A A F P V F A R I H Q Q V W D L
CTCGATGTGCCCCGATCTGGAGGTGAACGAGACCGGTTACGCCAGCCGGC 2200
15 L D V P D L E V N E T G Y A Q P A
CCTGTTTCGAATGCAGGTGGCTCTGTTCCGGGCTGCTGGAATCGTGGGGTG 2250
L F A M Q V A L F G L L E S W G
TACGACCGGACGCGGTGATCGGCCATTGCGTGGGTGAGCTTGCGGCTGCG 2300
V R P D A V I G H S V G E L A A A
20 TATGTGTCCGGGGTGTGGTTCGTTGGAGGATGCCTGCACTTTGGTGTGCGC 2350
Y V S G V W S L E D A C T L V S A
GCGGGCTCGTCTGATGCAGGCTCTGCCCGCGGGTGGGGTGATGGTCGCTG 2400
R A R L M Q A L P A G G V M V A
TCCCGGTCTCGGAGGATGAGGCCCCGGGCGTCTGGGTGAGGGTGTGGAG 2450
25 V P V S E D E A R A V L G E G V E
ATCGCCGCGGTCAACGGCCCGTCTCGGTGGTTCTCTCCGGTGATGAGGC 2500
I A A V N G P S S V V L S G D E A
CGCCGTGCTGCAGGCCGCGGAGGGGCTGGGGAAGTGGACGCGGCTGGCGA 2550
A V L Q A A E G L G K W T R L A
30 CCAGCCACGCGTTCATTCCGCCCCGATGGAACCCATGCTGGAGGAGTTC 2600
T S H A F H S A R M E P M L E E F
CGGGCGGTGCGCGAAGGCCTGACCTACCGGACGCGCAGGTCTCCATGGC 2650
R A V A E G L T Y R T P Q V S M A
CGTTGGTGATCAGGTGACCACCGCTGAGTACTGGGTGCGGCAGGTCCGGG 2700
35 V G D Q V T T A E Y W V R Q V R
ACACGGTCCGGTTCGGCGAGCAGGTGGCCTCGTACGAGGACGCCGTGTTC 2750
D T V R F G E Q V A S Y E D A V F
GTCGAGCTGGGTGCCGACCGGTCACTGGCCCCGCTGGTTCGACGGTGTGCG 2800
V E L G A D R S L A R L V D G V A
40 GATGTCACGCGCGACACGAAATCCAGGCCGCGATCGGCGCCCTGGCCC 2850
M L H G D H E I Q A A I G A L A
ACCTGTATGTCAACGGCGTCAACGGTCGACTGGCCCCGCTCCTGGGCGAT 2900
H L Y V N G V T V D W P A L L G D
GCTCCGGCAACACGGGTGCTGGACCTTCCGACATACGCTTCCAGCACCA 2950
45 A P A T R V L D L P T Y A F Q H Q
GCGCTACTGGCTCGAGTCGGCTCCCCCGGCCACGGCCGACTCGGGCCACC 3000
R Y W L E S A P P A T A D S G H
CCGTCTCGGCACCGGAGTCGCGTCCGCGGGTCCGCGGGCCGGGTGTTT 3050
P V L G T G V A V A G S P G R V F
50 ACGGGTCCCGTGGCCCGGGTGGCGACCGCGCGGTGTTTCATCGCCGAAC 3100
T G P V P A G A D R A V F I A E L
GGCGCTCGCCGCCCGGACGCCACCGACTGCGCCACGGTCGAACAGCTCG 3150
A L A A A D A T D C A T V E Q L
ACGTACCTCCGTGCCCCGGCGGATCCGCCCCGCGCAGGGCCACCGCGCAG 3200

- 119 -

D V T S V P G G S A R G R A T A Q
ACCTGGGTGATGAACCCGCGCCGACGGGCGGCGCGCTTCACCGTCCA 3250
T W V D E P A A D G R R R F T V H
CACCCGCGTCGGCGACGCCCCGTGGACGCTGCACGCCGAGGGGGTTCTCC 3300
5 T R V G D A P W T L H A E G V L
GCCCCGCGCGTGTCCCGAGCCCGAAGCCGTGACACCGCCTGGCCCCCG 3350
R P G R V P Q P E A V D T A W P P
CCGGGCGCGGTGCCCCGCGACGGGCTGCCCCGGGCGTGGCGACGCGCGGA 3400
P G A V P A D G L P G A W R R A D
10 CCAGGTCTTCGTGAAGCCGAAGTCGACAGCCCTGACGGCTTCGTGGCAC 3450
Q V F V E A E V D S P D G F V A
ACCCCGACCTGCTCGACGCGGTCTTCTCCGCGGTGCGCGACGGGAGCCGC 3500
H P D L L D A V F S A V G D G S R
CAGCCGACCGGATGGCGCGACCTCGCGGTGCACGCGTCGGACGCCACCGT 3550
15 Q P T G W R D L A V H A S D A T V
GCTGCGCGCTGCCTCACCCGCCGCGACAGTGGTGTGCTGGAGCTCGCCG 3600
L R A C L T R R D S G V V E L A
CCTTCGACGGTGCCGGAATGCCGGTGCTCACCGCGGAGTCGGTGACGCTG 3650
A F D G A G M P V L T A E S V T L
20 GGCGAGGTGCGCTCGGCAGGCGGATCCGACGAGTCGGACGGTCTGCTTCG 3700
G E V A S A G G S D E S D G L L R
GCTTGAGTGGTTGCCGGTGGCGGAGGCCCCACTACGACGGTGCCGACGAGC 3750
L E W L P V A E A H Y D G A D E
TGCCCGAGGGCTACACCCTCATCACGCCACACACCCCGACGACCCCGAC 3800
25 L P E G Y T L I T A T H P D D P D
GACCCCAACCAACCCCAACAACACACCCACACGACCCACACACAAACCAC 3850
D P T N P H N T P T R T H T Q T T
ACGCGTCTCACCGCCCTCCAACACCACCTCATCACCAACCAACCAACCC 3900
R V L T A L Q H H L I T T N H T
30 TCATCGTCCACACCACCACCGACCCCCAGGCGCGCGCGTCACCGGCCTC 3950
L I V H T T T D P P G A A V T G L
ACCCGCACCGCACAAAACGAACACCCCGCGCATCCACCTCATCGAAAC 4000
T R T A Q N E H P G R I H L I E T
CCACCACCCCAACACCCCACTCCCCCTCACCCAACCTACCACCCCTCCACC 4050
35 H H P H T P L P L T Q L T T L H
AACCCCACTACGCCTACCAACAACACCCTCCACACCCCACTCACC 4100
Q P H L R L T N N T L H T P H L T
CCCATCACCAACCAACAACACCACCAACCAACCCCAACACCCCAAC 4150
P I T T H H N T T T T T P N T P P
40 CCTCAACCCCAACCAACCATCCTCATACCGGCGGCTCCGGCACCCCTCG 4200
L N P N H A I L I T G G S G T L
CCGGCATCCTCGCCCGCCACCTCAACCAACCCCAACCTACCTCCTCTCC 4250
A G I L A R H L N H P H T Y L L S
CGCACACCAACCAACCCCAACCAACCCCGCACCCACATCCCCTGCGACCT 4300
45 R T P P P P T T P G T H I P C D L
CACCGACCCCAACCAAAATCACCAAGCCCTCACCCACATACCACAACCCC 4350
T D P T Q I T Q A L T H I P Q P
TCACCGGCATCTTCACACCGCGCCACCCTCGACGACGCCACCCTCACC 4400
L T G I F H T A A T L D D A T L T
50 AACCTCACCCCAACACCTCACCACCACCCTCCAACCCAAAGCCGACGC 4450
N L T P Q H L T T T L Q P K A D A
CGCCTGGCACCTCCACCACCAACCCAAACCAACCCCTCACCCACTTCG 4500
A W H L H H H T Q N Q P L T H F
TCCTCTACTCCAGCGCGCGCCACCCCTCGGCAGCCCCGGCCAAGCCAAC 4550

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- 120 -

V L Y S S A A A T L G S P G Q A N
TACGCCGCCGCAACGCCTTCCTCGACGCCCTCGCCACCCACCGCCACAC 4600
Y A A A N A F L D A L A T H R H T
CCAAGGACAACCCGCCACCACCATCGCCTGGGGCATGTGGCACACCACCA 4650
5 Q G Q P A T T I A W G M W H T T
CCACACTCACCAGCCAACCTACCGACAGCGACCGCGACCGCATCCGCCGC 4700
T T L T S Q L T D S D R D R I R R
GGCGGCTTCCTGCCGATCTCGGACGACGAGGGCATGC
10 G G F L P I S D D E G M

The *NheI-XhoI* hybrid FK-506 PKS module 8 containing the AT domain of
module 13 of rapamycin is shown below.

GCATGCGGCTGTACGAGGCGGCACGGCGCACCGGAAGTCCCGTGGTGGTG 50
M R L Y E A A R R T G S P V V V
15 GCGGCCGCGCTCGACGACGCGCCGACGTGCCGCTGCTGCGCGGGCTGCG 100
A A A L D D A P D V P L L R G L R
GCGTACGACCGTCCGGCGTGCCGCCGTCCGGGAACGCTCTCTCGCCGACC 150
R T T V R R A A V R E R S L A D
GCTCGCCGTGCTGCCCCGACGACGAGCGCGCCGACGCCTCCCTCGCGTTG 200
20 R S P C C P T T S A P T P P S R S
TCCTGGAACAGCACCGCCACCGTGCTCGGCCACCTGGGCGCCGAAGACAT 250
S W N S T A T V L G H L G A E D I
CCCGGCGACGACGACGTTCAAGGAACCTCGGCATCGACTCGCTCACC GCG 300
P A T T T F K E L G I D S L T A
25 TCCAGCTGCGCAACGCGCTGACCACGGCGACCGGCGTACGCCTCAACGCC 350
V Q L R N A L T T A T G V R L N A
ACAGCGGTCTTCGACTTTCCGACGCGCGCGCGCTCGCCGCGAGACTCGG 400
T A V F D F P T P R A L A A R L G
CGACGAGCTGGCCGGTACCCGCGCGCCCGTCCGCGCCCGGACCGCGGCCA 450
30 D E L A G T R A P V A A R T A A
CCGCGGCCGCGCACGACGAACCGCTGGCGATCGTGGGCATGGCCTGCCGT 500
T A A A H D E P L A I V G M A C R
CTGCCGGGCGGGTTCGCGTCGCCACAGGAGCTGTGGCGTCTCGTCGCGTC 550
L P G G V A S P Q E L W R L V A S
35 CGGCACCGACGCCATCACGGAGTTCCCCGCGGACCGCGGCTGGGACGTGG 600
G T D A I T E F P A D R G W D V
ACGCGCTCTACGACCCGGACCCCGACGCGATCGGCAAGACCTTCGTCCGG 650
D A L Y D P D P D A I G K T F V R
CACGGCGGCTTCCTCGACGGTGCGACCGGCTTCGACGCGGCGTTCTTCGG 700
40 H G G F L D G A T G F D A A F F G
GATCAGCCCGCGGAGGCCCTGGCCATGGACCCGACGCAACGGGTGCTCC 750
I S P R E A L A M D P Q Q R V L
TGGAGACGTCTGGGAGGCGTTCGAAAGCGCGGGCATCACCCCGGACGCG 800
L E T S W E A F E S A G I T P D A
45 GCGCGGGGACGCGACACCGGCGTGTTCATCGGCGCGTTCCTACGGGTA 850
A R G S D T G V F I G A F S Y G Y
CGGCACGGGTGCGGATACCAACGGCTTCGGCGCGACAGGGTCGACAGACCA 900
G T G A D T N G F G A T G S Q T
CGGTGCTCTCCGGCCGCTCTCGTACTTCTACGGTCTGGAGGGCCCTTCG 950
50 S V L S G R L S Y F Y G L E G P S
GTCACGGTGCACACCGCCTGCTCGTCTGCTCACTGGTTCGCCCTGCACCAGGC 1000
V T V D T A C S S S L V A L H Q A

- 121 -

AGGGCAGTCCCTGCGCTCGGGCGAATGCTCGCTCGCCCTGGTCGGCGGTG 1050
G Q S L R S G E C S L A L V G G
TCACGGTGATGGCGTCGCCCCGGCGGATTCTCGAGTTCTCCCGGCAGCGC 1100
V T V M A S P G G F V E F S R Q R
5 GGGCTCGCGCCGACGGGCGGGCGAAGGCGTTTCGGCGCGGGCGCGGACGG 1150
G L A P D G R A K A F G A G A D G
TACGAGCTTCGCGGAGGGCGCCGGTGCCCTGGTGGTCGAGCGGCTCTCCG 1200
T S F A E G A G A L V V E R L S
ACGCGGAGCGCCACGGCCACACCGTCTCGCCCTCGTACGCGGCTCCGCG 1250
10 D A E R H G H T V L A L V R G S A
GCTAACTCCGACGGCGCGTCGAACGGTCTGTTCGGCGCCGAACGGCCCCCTC 1300
A N S D G A S N G L S A P N G P S
CCAGGAACGCGTCATCCACCAGGCCCTCGCGAACGCGAAACTCACCCCCG 1350
Q E R V I H Q A L A N A K L T P
15 CCGATGTCGACGCGGTTCGAGGCGCACGGCACCGGCACCCGCCTCGGCGAC 1400
A D V D A V E A H G T G T R L G D
CCCATCGAGGCGCAGGCGCTGCTCGCGACGTACGGACAGGACCGGGCGAC 1450
P I E A Q A L L A T Y G Q D R A T
20 GCCCTGCTGCTCGGCTCGCTGAAGTCGAACATCGGGCACGCCAGGCCG 1500
P L L L G S L K S N I G H A Q A
CGTCAGGGTTCGCGGGATCATCAAGATGGTGCAGGCCATCCGGCACGGG 1550
A S G V A G I I K M V Q A I R H G
GAACTGCCGCCGACACTGCACGCGGACGAGCCGTGCGCGCACGTGACTG 1600
E L P P T L H A D E P S P H V D W
25 GACGGCCCGGTGCCGTCGAGCTCCTGACGTGCGCCCGGCCGTGGCCGGGGA 1650
T A G A V E L L T S A R P W P G
CCGGTCGCCCCGCGCCGCTGCCGTCTCGTCGTTTCGGCGTGAGCGGCACG 1700
T G R P R R A A V S S F G V S G T
30 AACGCCACATCATCTTGAGGCAGGACCGGTCAAACGGGACCGGTCTGA 1750
N A H I I L E A G P V K T G P V E
GGCAGGAGCGATCGAGGCAGGACCGGTCTGAAGTAGGACCGGTCTGAGGCTG 1800
A G A I E A G P V E V G P V E A
GACCGTCCCCCGCGGCGCCGCCGTCTCAGCACCGGGCGAAGACCTTCCGCTG 1850
G P L P A A P P S A P G E D L P L
35 CTCGTGTCGGCGCGTTCCCCCGAGGCACTCGACGAGCAGATCGGGCGCCT 1900
L V S A R S P E A L D E Q I G R L
GCGCGCTATCTCGACACCGGCCCCGGCGTCGACCGGGCGGCCGTGGCGC 1950
R A Y L D T G P G V D R A A V A
AGACACTGGCCCCGGGTACGCACTTCACCCACCGGGCCGTACTGCTCGGG 2000
40 Q T L A R R T H F T H R A V L L G
GACACCGTCATCGGCGCTCCCCCGCGGACCAGGCCGACGAACCTGTCTT 2050
D T V I G A P P A D Q A D E L V F
CGTCTACTCCGGTCAGGGCACCCAGCATCCCGCGATGGGCGAGCAGCTAG 2100
V Y S G Q G T Q H P A M G E Q L
45 CCGATTCTGTCGGTGGTGTTCGCGGAGCGGATGGCCGAGTGTGCGGCGGCG 2150
A D S S V V F A E R M A E C A A A
TTGCGCGAGTTCGTGGACTGGGATCTGTTACGGTTCTGGATGATCCGGC 2200
L R E F V D W D L F T V L D D P A
GGTGGTGGACCGGTTGATGTGGTCCAGCCCGCTTCTGGGCGATGATGG 2250
50 V V D R V D V V Q P A S W A M M
TTTCCCTGGCCGCGGTGTGGCAGGCGGCCGGTGTGCGGCCGGATGCGGTG 2300
V S L A A V W Q A A G V R P D A V
ATCGGCCATTTCGAGGGTGAGATCGCCGCAGCTTGTGTGGCGGGTGCGGT 2350
I G H S Q G E I A A A C V A G A V

- 122 -

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GTCACTACGCGATGCCGCCCGGATCGTGACCTTGCGCAGCCAGGCGATCG 2400
S L R D A A R I V T L R S Q A I
CCCCGGGCGCTGGCGGGCGGGGCGCGATGGCATCCGTCGCCCTGCCCGCG 2450
A R G L A G R G A M A S V A L P A
CAGGATGTCGAGCTGGTCGACGGGGCCTGGATCGCCGCCACAACGGGCC 2500
Q D V E L V D G A W I A A H N G P
CGCCTCCACCGTGATCGCGGGCACCCCGGAAGCGGTGACCATGTCCTCA 2550
A S T V I A G T P E A V D H V L
CCGCTCATGAGGCACAAGGGGTGCGGGTGCGGCGGATCACCGTCGACTAT 2600
T A H E A Q G V R V R R I T V D Y
GCCTCGCACACCCCGCACGTCGAGCTGATCCGCGACGAACACTACTCGACAT 2650
A S H T P H V E L I R D E L L D I
CACTAGCGACAGCAGCTCGCAGACCCCGCTCGTGCCGTGGCTGTGACCG 2700
T S D S S S Q T P L V P W L S T
TGGACGGCACCTGGGTCGACAGCCCGCTGGACGGGGAGTACTGGTACCGG 2750
V D G T W V D S P L D G E Y W Y R
AACCTGCGTGAACCGGTGCGTTTCCACCCCGCGTCAGCCAGTTGCAGGC 2800
N L R E P V G F H P A V S Q L Q A
CCAGGGCGACACCGTGTTGTCGAGGTGAGCGCCAGCCCGGTGTTGTTGC 2850
Q G D T V F V E V S A S P V L L
AGGCGATGGACGATGTCGTCACGGTTGCCACGCTGCGTCGTGACGAC 2900
Q A M D D D V V T V A T L R R D D
GGCGACGCCACCCGGATGCTCACCGCCCTGGCACAGGCCTATGTCCACGG 2950
G D A T R M L T A L A Q A Y V H G
CGTCACCGTCGACTGGCCCCGCATCCTCGGCACCACCACAACCCGGGTAC 3000
V T V D W P A I L G T T T T R V
TGGACCTTCGACCTACGCCTTCCAACACCAGCGGTACTGGCTCGAGTCG 3050
L D L P T Y A F Q H Q R Y W L E S
GCTCCCCCGGCCACGGCCGACTCGGGCCACCCCGTCTCGGCACCGGAGT 3100
A P P A T A D S G H P V L G T G V
CGCCGTGCGCGGGTGC CGGGCCGGGTGTTACGGGTCCCGTGCCCGCCG 3150
A V A G S P G R V F T G P V P A
GTGCGGACCGCGCGGTGTTTCATCGCCGAACTGGCGCTCGCCGCCGCCGAC 3200
G A D R A V F I A E L A L A A A D
GCCACCGACTGCGCCACGGTGAACAGCTCGACGTACCTCCGTGCCCGG 3250
A T D C A T V E Q L D V T S V P G
CGGATCCGCCCCGGGCAGGGCCACCGCGCAGACCTGGGTGATGAACCCG 3300
G S A R G R A T A Q T W V D E P
CCGCGACGGGCGGCGCGCTTACCGTCCACACCCGCGTCGGCGACGCC 3350
A A D G R R R F T V H T R V G D A
CCGTGGACGCTGCACGCCGAGGGGTTCTCCGCCCCGGCCGCGTGCCCA 3400
P W T L H A E G V L R P G R V P Q
GCCCCAAGCCGTGACACCGCCTGGCCCCCGCGGGCGCGGTGCCCGCGG 3450
P E A V D T A W P P P G A V P A
ACGGGCTGCCCCGGGCGTGGCGACGCGCGGACCAGGTCTTCGTGCAAGCC 3500
D G L P G A W R R A D Q V F V E A
GAAGTCGACAGCCCTGACGGCTTCGTGGCACACCCGACCTGCTCGACGC 3550
E V D S P D G F V A H P D L L D A
GGTCTTCTCCGCGGTGCGCGACGGGAGCCGCCAGCCGACCGGATGGCGCG 3600
V F S A V G D G S R Q P T G W R
ACCTCGCGGTGCACGCGTCGGACGCCACCGTGCTGCGCGCCTGCCTCACC 3650
D L A V H A S D A T V L R A C L T
CGCCGCGACAGTGGTGTCGTGGAGCTCGCCGCTTCGACGGTGCCGGAAT 3700
R R D S G V V E L A A F D G A G M

- 123 -

GCCGGTGCTCACC GCGGAGTCGGTGACGCTGGGCGAGGTCGCGTCGGCAG 3750
P V L T A E S V T L G E V A S A
GCGGATCCGACGAGTCGGACGGTCTGCTTCGGCTTGAGTGGTTGCCGGTG 3800
G G S D E S D G L L R L E W L P V
5 GCGGAGGCCCCACTACGACGGTGCCGACGAGCTGCCCCAGGGCTACACCCT 3850
A E A H Y D G A D E L P E G Y T L
CATCACCGCCACACACCCCCGACGACCCCGACGACCCACCAACCCCCACA 3900
I T A T H P D D P D D P T N P H
ACACACCCACACGACCCACACACAAACCACACGCGTCCTCACCGCCCTC 3950
10 N T P T R T H T Q T T R V L T A L
CAACACCACCTCATCACCACCAACCACACCCTCATCGTCCACACCACCAC 4000
Q H H L I T T N H T L I V H T T T
CGACCCCCCAGGCGCCGCGGTACCGGCCTACCCGACCGCACAAAACG 4050
D P P G A A V T G L T R T A Q N
15 AACACCCCGGCGCATCCACCTCATCGAAACCCACCACCCCAACCCCA 4100
E H P G R I H L I E T H H P H T P
CTCCCCCTCACCAACTCACCACCTCCACCAACCCCACTACGCCTCAC 4150
L P L T Q L T T L H Q P H L R L T
CAACAACACCTCCACACCCCCACCTACCCCCATCACCACCCACCACA 4200
20 N N T L H T P H L T P I T T H H
ACACCACCAACCAACCCCAACACCCACCCCTCAACCCCAACCACGCC 4250
N T T T T T P N T P P L N P N H A
ATCCTCATCACCGGCGGCTCCGGCACCCCTCGCCGGCATCTCGCCGCCA 4300
I L I T G G S G T L A G I L A R H
25 CCTCAACCAACCCCAACCTACCTCCTCCTCCCGCACACCACCCCA 4350
L N H P H T Y L L S R T P P P P
CCACACCCGGCACCCACATCCCCTGCGACCTACCGACCCCAACCAATC 4400
T T P G T H I P C D L T D P T Q I
ACCCAAGCCCTCACCCACATAACCACAACCCCTACCGGCATCTTCCACAC 4450
30 T Q A L T H I P Q P L T G I F H T
CGCCGCCACCCCTCGACGACGCCACCCCTACCAACCTACCCCCCAACACC 4500
A A T L D D A T L T N L T P Q H
TCACCACCACCTCCAACCCAAAGCCGACGCCGCTGGCACCTCCACCAC 4550
L T T T L Q P K A D A A W H L H H
35 CACACCCAAAACCAACCCCTCACCCACTTCGTCTCTACTCCAGCGCCGC 4600
H T Q N Q P L T H F V L Y S S A A
CGCCACCCCTCGGCAGCCCCGGCCAAGCCAACTACGCCGCCGCCAACGCCT 4650
A T L G S P G Q A N Y A A A N A
TCCTCGACGCCCTCGCCACCCACCGCCACCCCAAGGACAACCCGCCACC 4700
40 F L D A L A T H R H T Q G Q P A T
ACCATCGCCTGGGGCATGTGGCACACCACCACTCACCAGCCAACT 4750
T I A W G M W H T T T T L T S Q L
CACCGACAGCGACCGCGACCGCATCCGCCGCGGCGCTTCTGCCGATCT 4800
T D S D R D R I R R G G F L P I
45 CGGACGACGAGGGCATGC
S D D E G M

Example 3

Recombinant PKS Genes for 13-desmethoxy FK-506 and FK-520

50 The present invention provides a variety of recombinant PKS genes in addition to those described in Examples 1 and 2 for producing 13-desmethoxy FK-506 and FK-520

- 124 -

compounds. This Example provides the construction protocols for recombinant FK-520 and FK-506 (from *Streptomyces* sp. MA6858 (ATCC 55098), described in U.S. Patent Nos. 5,116,756, incorporated herein by reference) PKS genes in which the module 8 AT coding sequences have been replaced by either the *rapAT3* (the AT domain from module 3 of the rapamycin PKS), *rapAT12*, *eryAT1* (the AT domain from module 1 of the erythromycin (DEBS) PKS), or *eryAT2* coding sequences. Each of these constructs provides a PKS that produces the 13-desmethoxy-13-methyl derivative, except for the *rapAT12* replacement, which provides the 13-desmethoxy derivative, i.e., it has a hydrogen where the other derivatives have methyl.

Figure 7 shows the process used to generate the AT replacement constructs. First, a fragment of ~4.5 kb containing module 8 coding sequences from the FK-520 cluster of ATCC 14891 was cloned using the convenient restriction sites *SacI* and *SphI* (Step A in Figure 7). The choice of restriction sites used to clone a 4.0 - 4.5 kb fragment comprising module 8 coding sequences from other FK-520 or FK-506 clusters can be different depending on the DNA sequence, but the overall scheme is identical. The unique *SacI* and *SphI* restriction sites at the ends of the FK-520 module 8 fragment were then changed to unique *Bgl* II and *Nsi* I sites by ligation to synthetic linkers (described in the preceding Examples, see Step B of Figure 7). Fragments containing sequences 5' and 3' of the AT8 sequences were then amplified using primers, described above, that introduced either an *Avr* II site or an *Nhe* I site at two different KS/AT boundaries and an *Xho* I site at the AT/DH boundary (Step C of Figure 7). Heterologous AT domains from the rapamycin and erythromycin gene clusters were amplified using primers, as described above, that introduced the same sites as just described (Step D of Figure 7). The fragments were ligated to give hybrid modules with in-frame fusions at the KS/AT and AT/DH boundaries (Step E of Figure 7). Finally, these hybrid modules were ligated into the *Bam* HI and *Pst* I sites of the KC515 vector. The resulting recombinant phage were used to transform the FK-506 and FK-520 producer strains to yield the desired recombinant cells, as described in the preceding Examples.

- 125 -

The following table shows the location and sequences surrounding the engineered site of each of the heterologous AT domains employed. The FK-506 hybrid construct was used as a control for the FK-520 recombinant cells produced, and a similar FK-520 hybrid construct was used as a control for the FK-506 recombinant cells.

5

Heterologous AT	Enzyme	Location of Engineered Site
FK-506 AT8 (hydroxymalonyl)	<i>AvrII</i>	GGCCGT <u>ccgcgc</u> CGTGCGGCGGTCTCGTCGTTTC G R P R R A A V S S F
	<i>NheI</i>	ACCCAGCATCCCGCGATGGGTGAGCG <u>gctcgc</u> C T Q H P A M G E R L A
	<i>XhoI</i>	TACGCCTTCCAGCGGCGGCCCTACTGG <u>atcgag</u> Y A F Q R R P Y W I E
rapamycin AT3 (methylmalonyl)	<i>AvrII</i>	GACCGG <u>ccccgt</u> CGGGCGGGCGTGTCGTCTTC D R P R R A G V S S F
	<i>NheI</i>	TGGCAGTGGCTGGGGATGGGCAGTGC <u>cctgcg</u> G W Q W L G M G S A L R
	<i>XhoI</i>	TACGCCTTCCAACACCAGCGGTACTGG <u>gtcgag</u> Y A F Q H Q R Y W V E
rapamycin AT12 (malonyl)	<i>AvrII</i>	GGCCGAg <u>cgcg</u> cCGGGCAGGCGTGTCGTCTTC G R A R R A G V S S F
	<i>NheI</i>	TCGCAGCGTGCTGGCATGGGTGAGGA <u>actggc</u> C S Q R A G M G E E L A
	<i>XhoI</i>	TACGCCTTCCAGCACCAGCGCTACTGG <u>ctcgag</u> Y A F Q H Q R Y W L E
DEBS AT1 (methylmalonyl)	<i>AvrII</i>	GCGCGA <u>ccgcgc</u> cCGGGCGGGGTCTCGTCGTTTC A R P R R A G V S S F
	<i>NheI</i>	TGGCAGTGGGCGGGCATGGCCGTGCA <u>octgct</u> C W Q W A G M A V D L L
	<i>XhoI</i>	TACCCGTTCCAGCGCGAGCGGTCTGG <u>ctcgaa</u> Y P F Q R E R V W L E
DEBS AT2 (methylmalonyl)	<i>AvrII</i>	GACGGG <u>gtgcgc</u> cCGGGCAGGTGTGTGCGGCGTTTC D G V R R A G V S A F
	<i>NheI</i>	GCCCAGTGGGAAGGCATGGCGCGGGA <u>gttggt</u> G A Q W E G M A R E L L
	<i>XhoI</i>	TATCCTTTCCAGGGCAAGCGTTCTGG <u>ctgctg</u> Y P F Q G K R F W L L

- 126 -

The sequences shown below provide the location of the KS/AT boundaries chosen in the FK-520 module 8 coding sequences. Regions where *AvrII* and *NheI* sites were engineered are indicated by lower case and underlining.

5 CCGGCGCCGTCGAACTGCTGACGTCGGCCCCGGCCGTGGCCCCGAGACCGACCGGccaacggc
A G A V E L L T S A R P W P E T D R P R
GTGCCCGCTCTCCTCGTTCGGGGTGAGCGGCACCAACGCCCACGTCATCCTGGAGGCCG
R A A V S S F G V S G T N A H V I L E A
GACCGGTAACGGAGAGCCCCGCGGCATCGCCTTCCGGTGACCTTCCCCTGCTGGTGTGCG
10 G P V T E T P A A S P S G D L P L L V S
CACGCTCACCGGAAGCGCTCGACGAGCAGATCCGCGACTGCGCGCTACCTGGACACCA
A R S P E A L D E Q I R R L R A Y L D T
CCCCGGACGTCGACCGGGTGGCCGTGGCACAGACGCTGGCCCCGGCGCACACACTTCGCC
T P D V D R V A V A Q T L A R R T H F A
ACCGCGCCGTGCTGCTCGGTGACACCGTCATCACACACCCCCCGGGACCGGCCCGGACG
15 H R A V L L G D T V I T T P P A D R P D
AACTCGTCTTCGTCTACTCCGGCCAGGGCACCCAGCATCCCGCGATGGGCGAGCAgctcg
E L V F V Y S G Q G T Q H P A M G E Q L
cCGCCGCCCATCCCGTGTTCGCCGACGCTGGCATGAAGCGCTCCGCCGCTTGACAACC
A A A H P V F A D A W H E A L R R L D N
20

The sequences shown below provide the location of the AT/DH boundary chosen in the FK-520 module 8 coding sequences. The region where an *XhoI* site was engineered is indicated by lower case and underlining.

25 TCCTCGGGGCTGGGTACGGCACGACGCGGATGTGCCCGGTACGCGTTCCAACGGCGGG
I L G A G S R H D A D V P A Y A F Q R R
ACTACTGGatcgagTCGGCACGCCCCGGCCGCATCCGACGCGGGCCACCCGTGCTGGGCT
H Y W I E S A R P A A S D A G H P V L G

The sequences shown below provide the location of the KS/AT boundaries chosen in the FK-506 module 8 coding sequences. Regions where *AvrII* and *NheI* sites were engineered are indicated by lower case and underlining.

30 TCGGCCAGGCCGTGGCCCGGACCGGCCGTcagcgcCGTGC GGCGGTCTCGTCTCGGTTCGGG
S A R P W P R T G R P R R A A V S S F G
GTGAGCGGCACCAACGCCCACATCATCCTGGAGGCCGACCCGACCGAGGAGGCCGTCG
35 V S G T N A H I I L E A G P D Q E E P S
GCAGAACCGGCCGGTGACCTCCCCTGCTCGTGTGCGCACGGTCCCCGGAGGCACTGGAC
A E P A G D L P L L V S A R S P E A L D
GAGCAGATCGGGCGCTGCGGACTATCTCGACGCGCCCCCGGCGTGGACCTGGCGGCC
E Q I G R L R D Y L D A A P G V D L A A
40 GTGGCGCGGACACTGGCCACGCGTACGCACTTCTCCCACCGCGCCGTACTGCTCGGTGAC
V A R T L A T R T H F S H R A V L L G D
ACCGTCATCACCGCTCCCCCGTGGAAACAGCCGGGCGAGCTCGTCTTCTGCTACTCGGGA
T V I T A P P V E Q P G E L V F V Y S G
45 CAGGGCACCCAGCATCCCGCGATGGGTGAGCGgctcgcCGCAGCCTTCCCCGTGTTGCC
Q G T Q H P A M G E R L A A A F P V F A

- 127 -

GACCCGGACGTACCCGCCTACGCCTTCCAGCGGGCGGCCCTACTGGATCGAGTCCGCGCCG
D P D V P A Y A F Q R R P Y W I E S A P

The sequences shown below provide the location of the AT/DH boundary chosen
5 in the FK-506 module 8 coding sequences. The region where an *Xho*I site was
engineered is indicated by lower case and underlining.

GACCCGGACGTACCCGCCTACGCCTTCCAGCGGGCGGCCCTACTGGatcgagTCCGCGCCG
D P D V P A Y A F Q R R P Y W I E S A P

10 Example 4

Replacement of Methoxyl with Hydrogen or Methyl at C-15 of FK-506 and FK-520

The methods and reagents of the present invention also provide novel FK-506 and
FK-520 derivatives in which the methoxy group at C-15 is replaced by a hydrogen or
methyl. These derivatives are produced in recombinant host cells of the invention that
15 express recombinant PKS enzymes the produce the derivatives. These recombinant PKS
enzymes are prepared in accordance with the methodology of Examples 1 and 2, with the
exception that AT domain of module 7, instead of module 8, is replaced. Moreover, the
present invention provides recombinant PKS enzymes in which the AT domains of both
modules 7 and 8 have been changed. The table below summarizes the various compounds
20 provided by the present invention.

Compound	C-13	C-15	Derivative Provided
FK-506	hydrogen	hydrogen	13, 15-didesmethoxy-FK-506
FK-506	hydrogen	methoxy	13-desmethoxy-FK-506
25 FK-506	hydrogen	methyl	13,15-didesmethoxy-15-methyl-FK-506
FK-506	methoxy	hydrogen	15-desmethoxy-FK-506
FK-506	methoxy	methoxy	Original Compound -- FK-506
FK-506	methoxy	methyl	15-desmethoxy-15-methyl-FK-506
FK-506	methyl	hydrogen	13,15-didesmethoxy-13-methyl-FK-506
30 FK-506	methyl	methoxy	13-desmethoxy-13-methyl-FK-506
FK-506	methyl	methyl	13,15-didesmethoxy-13,15-dimethyl-FK-506
FK-520	hydrogen	hydrogen	13, 15-didesmethoxy FK-520

- 128 -

FK-520	hydrogen	methoxy	13-desmethoxy FK-520
FK-520	hydrogen	methyl	13,15-didesmethoxy-15-methyl-FK-520
FK-520	methoxy	hydrogen	15-desmethoxy-FK-520
FK-520	methoxy	methoxy	Original Compound -- FK-520
5 FK-520	methoxy	methyl	15-desmethoxy-15-methyl-FK-520
FK-520	methyl	hydrogen	13,15-didesmethoxy-13-methyl-FK-520
FK-520	methyl	methoxy	13-desmethoxy-13-methyl-FK-520
FK-520	methyl	methyl	13,15-didesmethoxy-13,15-dimethyl-FK-520

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Example 5

Replacement of Methoxyl with Ethyl at C-13 and/or C-15 of FK-506 and FK-520

The present invention also provides novel FK-506 and FK-520 derivative compounds in which the methoxy groups at either or both the C-13 and C-15 positions are instead ethyl groups. These compounds are produced by novel PKS enzymes of the invention in which the AT domains of modules 8 and/or 7 are converted to ethylmalonyl specific AT domains by modification of the PKS gene that encodes the module.

Ethylmalonyl specific AT domain coding sequences can be obtained from, for example, the FK-520 PKS genes, the niddamycin PKS genes, and the tylosin PKS genes. The novel PKS genes of the invention include not only those in which either or both of the

AT domains of modules 7 and 8 have been converted to ethylmalonyl specific AT domains but also those in which one of the modules is converted to an ethylmalonyl specific AT domain and the other is converted to a malonyl specific or a methylmalonyl specific AT domain.

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Example 6

Neurotrophic Compounds

The compounds described in Examples 1 - 4, inclusive have immunosuppressant activity and can be employed as immunosuppressants in a manner and in formulations similar to those employed for FK-506. The compounds of the invention are generally effective for the prevention of organ rejection in patients receiving organ transplants and

- 129 -

in particular can be used for immunosuppression following orthotopic liver transplantation. These compounds also have pharmacokinetic properties and metabolism that are more advantageous for certain applications relative to those of FK-506 or FK-520. These compounds are also neurotrophic; however, for use as neurotrophins, it is desirable to modify the compounds to diminish or abolish their immunosuppressant activity. This can be readily accomplished by hydroxylating the compounds at the C-18 position using established chemical methodology or novel FK-520 PKS genes provided by the present invention.

Thus, in one aspect, the present invention provides a method for stimulating nerve growth that comprises administering a therapeutically effective dose of 18-hydroxy-FK-520. In another embodiment, the compound administered is a C-18,20-dihydroxy-FK-520 derivative. In another embodiment, the compound administered is a C-13-desmethoxy and/or C-15-desmethoxy 18-hydroxy-FK-520 derivative. In another embodiment, the compound administered is a C-13-desmethoxy and/or C-15-desmethoxy 18,20-dihydroxy-FK-520 derivative. In other embodiments, the compounds are the corresponding analogs of FK-506. The 18-hydroxy compounds of the invention can be prepared chemically, as described in U.S. Patent No. 5,189,042, incorporated herein by reference, or by fermentation of a recombinant host cell provided by the present invention that expresses a recombinant PKS in which the module 5 DH domain has been deleted or rendered non-functional.

The chemical methodology is as follows. A compound of the invention (~200 mg) is dissolved in 3 mL of dry methylene chloride and added to 45 μ L of 2,6-lutidine, and the mixture stirred at room temperature. After 10 minutes, tert-butyldimethylsilyl trifluoromethanesulfonate (64 μ L) is added by syringe. After 15 minutes, the reaction mixture is diluted with ethyl acetate, washed with saturated bicarbonate, washed with brine, and the organic phase dried over magnesium sulfate. Removal of solvent *in vacuo* and flash chromatography on silica gel (ethyl acetate: hexane (1:2) plus 1% methanol) gives the protected compound, which is dissolved in 95% ethanol (2.2 mL) and to which is added 53 μ L of pyridine, followed by selenium dioxide (58 mg). The flask is fitted with a water condenser and heated to 70°C on a mantle. After 20 hours, the mixture is

- 130 -

cooled to room temperature, filtered through diatomaceous earth, and the filtrate poured into a saturated sodium bicarbonate solution. This is extracted with ethyl acetate, and the organic phase is washed with brine and dried over magnesium sulfate. The solution is concentrated and purified by flash chromatography on silica gel (ethyl acetate: hexane (1:2) plus 1% methanol) to give the protected 18-hydroxy compound. This compound is dissolved in acetonitrile and treated with aqueous HF to remove the protecting groups. After dilution with ethyl acetate, the mixture is washed with saturated bicarbonate and brine, dried over magnesium sulfate, filtered, and evaporated to yield the 18-hydroxy compound. Thus, the present invention provides the C-18-hydroxyl derivatives of the compounds described in Examples 1 - 4.

Those of skill in the art will recognize that other suitable chemical procedures can be used to prepare the novel 18-hydroxy compounds of the invention. See, e.g., Kawai *et al.*, Jan. 1993, Structure-activity profiles of macrolactam immunosuppressant FK-506 analogues, *FEBS Letters* 316(2): 107-113, incorporated herein by reference. These methods can be used to prepare both the C18-[S]-OH and C18-[R]-OH enantiomers, with the *R* enantiomer showing a somewhat lower IC₅₀, which may be preferred in some applications. See Kawai *et al.*, *supra*. Another preferred protocol is described in Umbreit and Sharpless, 1977, *JACS* 99(16): 1526-28, although it may be preferable to use 30 equivalents each of SeO₂ and t-BuOOH rather than the 0.02 and 3-4 equivalents, respectively, described in that reference.

All scientific and patent publications referenced herein are hereby incorporated by reference. The invention having now been described by way of written description and example, those of skill in the art will recognize that the invention can be practiced in a variety of embodiments, that the foregoing description and example is for purposes of illustration and not limitation of the following claims.